# SEARCH REQUEST FORM

# Scientific and Technical Information Center

Requester's Full Name: Margaref B Nadlan Examiner #: 60850 Date: 1-11-01  Art Unit: 17/4 Phone Number 308-35/B Serial Number: 09/549438 applif 200  Mail Box and Bldg/Room Location: 3-3 Results Format Preferred (circle): PAPER DISK E-MAIL  If m r than one s arch is submitt d, please prioritize searches in order f n ed.
Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched. Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or utility of the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc, if known. Please attach a copy of the cover sheet, pertinent claims, and abstract.  Examples of English Respective Concepts for the Sparation of English Respective Concepts (Please provide full names): De Planche, Thierry of all
Earliest Priority Filing Date: Secundary. The Compelo Afch. 10-Bond 18 *For Sequence Searches Only* Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the Security 6

STAFF USE ONLY Scarcher:	V	Vendors and cost where applicable
Searcher Phone #: X-4/39	AA Sequence (#)	
Searcher Location: E(C /700	Structure (#)	Questel/Orbit
Date Searcher Picked Up:		
ate Completed: 7-19-01	Litigation	
earcher Prep & Review Time:	· · · · · · · · · · · · · · · · · · ·	Sequence Systems
lerical Prep Time: /5	Patent Family	•
Online Time:	Other	

PTO-1590 (1-2000)

```
L32 80299 SEA FILE REGISTRY ABBEON PLUEON ?PHENYLALANIN?/CNS
L33 4672 SEA FILE REGISTRY ABBEON PLUEON ?NITRO?/CNS (L) L32
L34 757 SEA FILE REGISTRY ABBEON PLUEON N-4-NITRO? (L) L33
L35 1 SEA FILE REGISTRY ABBEON PLUEON 2-METHOXY? (L) L34
```

=> d 135

L35 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2001 ACS

RN 328406-65-1 REGISTRY

CN L-Phenylalanine, N-[(4-nitrophenoxy)carbonyl]-, 2-methoxyethyl ester

(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C19 H20 N2 O7

SR CA

=>

LC STN Files: CA, CAPLUS, TOXLIT

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

### => d all hitstr 137

```
ANSWER 1 OF 1 HCAPLUS
                            COPYRIGHT 2001 ACS
L37
AN
     2001:57848 HCAPLUS 4
```

DN 134:212851

TI Application of  $(S)-N_T/(4-n)$  itrophenoxycarbonyl) phenylalanine methoxyethyl ester as a new chiral derivatizing agent for proteinogenic amino acid analysis by high-performance liquid chromatography

事。 建二

ΑU Peter, A.; Vekes, E.; Torok, G.

- Department of Inorganic and Analytical Chemistry, University of Szeged, CS Szeged, 6720, Hung.
- Chromatographia (2000), 52(11/12), 821-826 SO CODEN: CHRGB7; ISSN: 0009-5893
- PB Friedrich Vieweg & Sohn Verlagsgesellschaft mbH
- DT
- LΆ English
- 64-3 (Pharmaceutical Analysis) CC
- The application of (S)-N-(4-nitrophenoxycarbonyl)phenylalanine AB methoxyethyl ester, | (S) NIFE, as a new chiral derivatizing agent for the resoln. of compds. possessing an amino group is described. Its applicability is demonstrated by the resoln. of proteinogenic amino acid enantiomers. The diastereomeric derivs, produced were sepd. by reversed-phase high-performance; liq. chromatog. The effects of pH, excess reagent and reaction time on the derivatization kinetics, and the effects of pH and the org. modifier on the sepn., were investigated.
- nitrophenoxycarbonyl phenylalanine deriv chiral agent HPLC; liq chromatog ST detn amino acid detn ·通信媒材(海山山市特点)。

IT

Reversed phase HPLC (sepn. of amino acids by reversed phase HPLC using (S)-N-(4nitrophenoxycarbonyl) phenylalanine methoxyethyl ester as chiral derivatizing agent)

52-90-4, L-Cysteine, analysis 56-41-7, L-Alanine, analysis IT 56-84-8, L-Aspartic acid, analysis 56-85-9, L-Serine, analysis L-Glutamine, analysis 56-86-0; L-Glutamic acid, analysis 56-87-1, L-Lysine, analysis 60-18-4, L-Tyrosine, analysis 61-90-5, L-Leucine, 63-68-3, L-Methionine, analysis 63-91-2, L-Phenylalanine, analysis 70-47-3, L-Asparagine, analysis 71-00-1, L-Histidine, 72-18-4, L-Valine, analysis 72-19-5, L-Threonine, analysis analysis 73-22-3, L-Tryptophan, analysis 73-32-3, L-Isoleucine, analysis 74-79-3, L-Arginine, analysis 147-85-3, L-Proline, analysis 302837-22÷5 1 302837+24-7 302837-19-0 302837-20-3 302837-25-8 302837-30+5 + 302837+31-6 302837-34-9 302837-27-0 302837-29-2 328406-66-2 328406-67-3 302837-35-0 302837-36-1 328406-68-4 328406-71-9 328406-72-0 328406-70-8 328406-69-5

RL: ANT (Analyte); ANST (Analytical study) (sepn. of amino acids by reversed phase HPLC using (S)-N-(4nitrophenoxycarbonyl) phenylalanine methoxyethyl ester as chiral derivatizing agent)

#### IT 328406-65-1

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (sepn. of amino acids by reversed phase HPLC using (S)-N-(4nitrophenoxycarbonyl) phenylalanine methoxyethyl ester as chiral derivatizing agent) 12-7 3:3 1 4 4 4

THE FIRE TO THE PARTY

,那都没有那一門的第二,前往1944年 \$2000 PAR 248 14 SE

RE.CNT 11

RE

6 79 76 1111 e, and by ...

- (1) Anon; J Biol Chem 1989, V264, P668 (学学)
- (2) Beesley, T; Chiral Chromatography 1998
- (3) Bojarski, J; Chem Anal 1997, V42, P139 HCAPLUS
- (4) Delplanche, T; Peptidomimetics Symposium 1999, 11
- (5) Gorog, S; Chromatogr B 1994, V659, P51 MEDLINE
- (6) Kleidernigg, O; Chromatographia 1997, V44, P465 HCAPLUS
- (7) Lunn, G; Handbook of Derivatization Reactions for HPLC 1998
- (8) Nimura, N; J Chromatogr 1980, V202, P375 HCAPLUS
- (9) Okamoto, Y; Chromatographic Enantiomer Separation on Chiral Polymers 1997

有金头蟹蹄一 化高键

网络萨拿好胡盒 电电流

TENERAL TENERA

- (10) Solvay; patent pending
- (11) Toyo'oka, T; Modern Derivatization Methods for Separation Science 1999
- IT 328406-65-1
  - RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (sepn. of amino acids by reversed phase HPLC using (S)-N-(4-nitrophenoxycarbonyl) phenylalanine methoxyethyl ester as chiral derivatizing agent)
- RN 328406-65-1 HCAPLUS
- CN L-Phenylalanine, N-[(4-nitrophénoxy)carbonyl]-, 2-methoxyethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L1 STR

```
NODE ATTRIBUTES:
HCOUNT
        IS M1
                    AT
                         10
NSPEC
         IS R
                    ΑT
                          2
NSPEC
         IS R
                    ΑT
         IS R
                    ΑT
NSPEC
NSPEC
         IS R
                    AT
NSPEC
         IS R
                    AT
                    AT
NSPEC
         IS R
                    AT
         IS C
NSPEC
         IS C
                          8
                    AT
NSPEC
         IS C
                    AT
NSPEC
         IS C
NSPEC
                    AT
                         10
NSPEC
         IS C
                    ΑT
                         11
NSPEC
         IS C
                    AT
                         12
         IS C
                    ΑT
                         13
NSPEC
         IS C
NSPEC
                    AT
```

DEFAULT MLEVEL IS ATOM MLEVEL IS CLASS AT 7 8 9 10 11 12 13 14

DEFAULT ECLEVEL IS LIMITED

## GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

L3	501	SEA FILE=REGISTRY SSS FUL L	1
L4	213	SEA FILE=HCAPLUS ABB=ON PL	U=ON L3
L6	1	SEA FILE=HCAPLUS ABB=ON PL	U=ON 2000:755247/AN
L14	75483	SEA FILE=HCAPLUS ABB=ON PL	U=ON (AMINO ACIDS OR PEPTIDES OR
	•	PROTEINS)/CC	
L15	83	SEA FILE=HCAPLUS ABB=ON PL	U=ON L4 AND L14
L16	82	SEA FILE=HCAPLUS ABB=ON PL	U=ON L15 NOT L6
L18	4	SEA FILE=HCAPLUS ABB=ON PL	U=ON (HPLC OR CHROMATOGARPH?) AND
		L16	
L19	9	SEA FILE=HCAPLUS ABB=ON PL	U=ON (ENANTIO? OR SEPARAT? OR
		CHROMOPHO? OR REAGENT) AND	L16
L20	12	SEA FILE=HCAPLUS ABB=ON PL	U=ON L18 OR L19
		•	

Timothy Saunders EIC-LAW Lib. 308-413

=>

```
COPYRIGHT 2001 ACS
     ANSWER 1 OF 12 HCAPLUS
L20
AN
     2001:220249 HCAPLUS
DN
     134:237834
TI
     Method for preparation of 2-[[N-(hydrocarbyloxycarbonyl)-L-
     alanyl]amino]thiazole-4-acetic acid ester derivatives
     Hirota, Yoshihiro; Matsunaga, Tomonori; Iwasaki, Fumiaki
IN
PA
     Tokuyama Corp., Japan
SO
     Jpn. Kokai Tokkyo Koho, 12 pp.
     CODEN: JKXXAF
     Patent
DΤ
     Japanese
LΑ
     ICM C07D277-44
IC
CC
     34-2 (Amino Acids, Peptides,
     Section cross-reference(s): 1, 26
FAN.CNT 1
     PATENT NO.
                       A2
PΙ
     JP 2001081083
OS
     CASREACT 134:237834; MARPAT 134:237834
```

The title compds. (I; R1 = C1-6 alkyl, C6-8 aryl, CH2Ph; R2 = C1-6 alkyl, AB Ph, CH2Ph) are prepd. by condensation of N-(hydrocarbyloxycarbonyl)-Lalanine (II; R1 = same as above) with 2-(2-amino-5-thiazolyl)-2methoxyiminoacetic acid ester (III; R2 = same as above) using a condensing agent, more specifically N, N'-carbonyldiimidazole and converted into 2-[2-[[N-(hydrocarbyloxycarbonyl)-L-alanyl]amino]thiazol-4-yl]-2methoxyiminoacetic acid I (R1 = same as above; R2 = H) by hydrolysis in the presence of base and neutralization. The latter 2-(2-aminothiazol-4yl)-2-methoxyiminoacețic acid derivs. are useful as intermediates for drugs, and in particular used as side chains for cephalosporin antibiotics. N, N'-carbonyldiimidazole is superior to other condensing agents such as dicyclohexylcarbodiimide and 1-ethyl-3-(3dimethylaminopropyl) carbodiimide hydrochloride and gives I in high yields. This process also gives III with high purity, enables the skipping of purifn. step for III prior to sapon., and simplifies and significantly improves the prodn. efficiency for the sapond. product, i.e. free acid I (R1 = same as above; R2 = H). Thus, 47.3 g N-tert-butoxycarbonyl-L-

GI

```
alanine and 57.3 g 2-(2-aminothiazol-4-yl)-2-(Z)-(methoxyimino) acetic acid
Et ester were added to 250 mL EtOAc, cooled to 0.degree., treated slowly
with 40.5 g N,N'-carbonyldiimidazole over a period of 10 min at
.ltoreq.5.degree., and stirred at 4.degree. for 1\ h. The reaction mixt.
was washed with 2 N HCl twice and 2 N NaOH and the org. layer was
sepd. to give, after evapn. of the solvent and silica gel
chromatog., 90.4 g I (R1 = tert-Bu, R2 = Et) (90.0% yield). In a sapon.
step, the org. layer obtained above was distd. in vacuo to the wt. of
113.2 g, treated with 67.2 g MeOH, and then slowly with 125 mL 3 N NaOH at
.ltoreq.25.degree. over a period of 15 min, and allowed to react at
25.degree. for 5 h. The solvent was distd. in vacuo from the reaction
mixt. until the mixt. weighed at 187.2 g, followed by adding 130 g H2O,
and the resulting mixt. was cooled 7.degree., treated slowly with 260 mL 2
N HCl over a period of 1 h, and stirred at .ltoreq.5.degree. for 2 h to
give, after centrifugation and washing the sepd. solid with
water and drying, 81.1 g I (R1 = tert-Bu, R2 = H) (87.0% yield).
alanylaminothiazolylmethoxyiminoacetic acid ester prepn intermediate
cephalosporin antibiotic; hydrocarbyloxycarbonylalanine condensation
aminothiazolylmethoxyiminoacetic acid ester; methoxyiminoacetic acid ester
alanyl aminothiazolyl prepn intermediate cephalosporin antibiotic
Condensation reaction
   (prepn. of [[N-(hydrocarbyloxycarbonyl)-L-alanyl]amino]thiazoleacetic
   acid ester derivs. by condensation of N-(hydrocarbyloxycarbonyl)-L-
   alanine with (aminothiazolyl) methoxyiminoacetic acid ester using
   N, N'-carbonyldiimidazole)
RL: PNU (Preparation, unclassified); PREP (Preparation)
   (.beta.-, antibiotics; prepn. of [[N-(hydrocarbyloxycarbonyl)-L-
   alanyl]amino]thiazoleacetic acid ester derivs. by condensation of
  N-(hydrocarbyloxycarbonyl)-L-alanine with (aminothiazolyl)methoxyiminoa
   cetic acid ester using N,N'-carbonyldiimidazole)
530-62-1, N,N'-Carbonyldiimidazole
                                    1142-20-7
                                                  15761-38-3
                                                               16639-86-4,
N-Ethoxycarbonyl-L-alanine
                             33294-53-0
                                          64485-88-7
                                                        65243-09-6
141695-27-4, N-Pheoxycarbonyl-L-alanine
                                          162281-00-7
                             oth Addition of
RL: RCT (Reactant)
   (prepn. of [[N-(hydrocarbyloxycarbonyl)-L-alanyl]amino]thiazoleacetic
   acid ester derivs. by condensation of N-(hydrocarbyloxycarbonyl)-L-
   alanine with (aminothiazolyl)methoxyiminoacetic acid ester using
                              N, N'-carbonyldiimidazole)
330566-03-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
   (prepn. of [[N-(hydrocarbyloxycarbonyl)-L-alanyl]amino]thiazoleacetic
   acid ester derivs. by condensation of N-(hydrocarbyloxycarbonyl)-L-
   alanine with (aminothiazolyl)methoxyiminoacetic acid ester using
                               Grand of the second
  N, N'-carbonyldiimidazole)
              330566-04-6P
                             330566-05-7P
                                            330566-06-8P 330566-07-9P
88970-81-4P
330566-08-0P
RL: SPN (Synthetic preparation); PREP (Preparation)
   (prepn. of [[N-(hydrocarbyloxycarbonyl)-L-alanyl]amino]thiazoleacetic
   acid ester derivs. by condensation of N-(hydrocarbyloxycarbonyl)-L-
   alanine with (aminothiazolyl) methoxyiminoacetic acid ester using
  N, N'-carbonyldiimidazole)
141695-27-4, N-Pheoxycarbonyl-L-alanine
RL: RCT (Reactant)
   (prepn. of [[N-(hydrocarbyloxycarbonyl)-L-alanyl]amino]thiazoleacetic
   acid ester derivs. by condensation of N-(hydrocarbyloxycarbonyl)-L-
   alanine with (aminothiazolyl) methoxyiminoacetic acid ester using
   N, N'-carbonyldiimidazole)
                               1966年21年1月1日 - 第二
141695-27-4 HCAPLUS
L-Alanine, N-(phenoxycarbonyl) - (9CI) (CA INDEX NAME)
```

The special sections

ST

IT

IT

IT

IT

IT

IT

RN

Absolute stereochemistry.

L20 ANSWER 2 OF 12 HCAPLUS COPYRIGHT 2001 ACS

AN 2000:853644 HCAPLUS

DN 134:193704

TI A solid-phase approach to analogues of the antibiotic mureidomycin

AU Bozzoli, Andrea; Kazmierski, Wieslaw; Kennedy, Gordon; Pasquarello, Alessandra; Pecunioso, Angelo

CS GlaxoWellcome SpA, Medicines Research Centre, Verona, 37135, Italy

SO Bioorg. Med. Chem. Lett. (2000), 10(24), 2759-2763

CODEN: BMCLE8; ISSN: 0960-894X

PB Elsevier Science Ltd.

DT Journal

LA English

CC 34-3 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 26, 33

GΙ

AB A library of 80 analogs of the antibacterial family of mureidomycins [(I); R = H, mureidomycin A; R = Gly, mureidomycin C], was prepd. using solid-phase chem. techniques. Analog fragments (2,3-diaminopropionic acid, methionine, p-tyrosine, m-tyrosine, as enantiomers, and tert-Bu 2-Ph malonate, as a racemate), were used in the synthesis. A selection of ten of the compds. with best a/a values (by LC-MS anal.) was presented. No inhibitory activity on the growth of S. aureus 853, E coli 1952, Saccharomyces cerevisiae NCY81, P. aeruginosa and P aeruginosa 2033 were detected for any library member.

ST mureidomycin analog combinatorial library prepn solid phase MSBAR

IT Combinatorial library

Solid phase synthesis

Structure-activity relationship

(prepn. of a combinatorial library of mureidomycin analogs using

```
; ,r:::12 cl
        solid-phase synthesis)
                                    Natural products
IT
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); BIOL (Biological study); PREP (Preparation)
        (prepn. of a combinatorial library of mureidomycin analogs using
                                    solid-phase synthesis)
     114797-04-5DP, Mureidomycin A, analogs
                                              327184-82-7P 327184-83-8P
IT
                                                   327184-87-2P
     327184-84-9P
                    327184-85-0P 327184-86-1P
                                                                  327184-88-3P
     327184-89-4P
                    327184-90-7P 327184-91-8P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); BIOL (Biological study); PREP (Preparation)
        (prepn. of a combinatorial library of mureidomycin analogs using
        solid-phase synthesis)
                               362-43-6 673-06-3, D-Phenylalanine
IT
     348-67-4, D-Methionine
     78342-42-4
                  109838-85-9
                                 198544-42-2 327184-80-5
     327184-93-0
     RL: RCT (Reactant)
        (prepn. of a combinatorial library of mureidomycin analogs using
        solid-phase synthesis)
                                  68691-77-0P 265321-20-8P
                   15083-09-7P
                                                                327184-50-9P
IT
     15083-05-3P
     327184-51-0P
                    327184-52-1P
                                    327184-53-2DP, resin-bound
                                                                  327184-53-2P
                                   327184-56-5P
                                                   327184-57-6P
     327184-54-3P
                    327184-55-4P
                                                                   327184-58-7P
                                    327184-61-2P 327184-62-3P
     327184-59-8P
                    327184-60-1P
                                                                   327184-63-4P
                                  327184-66-7P 327184-67-8P
     327184-64-5P
                    327184-65-6P
                                                                   327184-68-9P
     327184-69-0P 327184-70-3P
                                  327184-71+4DP, resin-bound
                   resin-bound 327184-73-6DP, resin-bound 327184-76-91 327184-78-1DP, resin-bound 327184-79-2DP, resin-bound
                                                                 327184-76-9DP,
     327184-72-5DP, resin-bound
     resin-bound
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of a combinatorial library) of mureidomycin analogs using solid-phase synthesis)
        solid-phase synthesis)
                    327184-75-8P 327184-77-0P 327184-81-6P
ΙT
     327184-74-7P
     RL: SPN (Synthetic preparation); PREP: (Preparation)
        (prepn. of a combinatorial library of mureidomycin analogs using
        solid-phase synthesis)
RE.CNT
                                   or Other William
RE
(1) Armstrong, A; Tetrahedron Lett 1988, V29, P2483 HCAPLUS
(2) Brandish, P; Antimicrob Agents Chemother 1996, V40, P1640 HCAPLUS
(3) Brandish, P; J Biol Chem 1996, V271, P7609 HCAPLUS
(4) Bugg, T; J Chem Soc, Perkin Trans 1 1999, P1279
(5) Bugg, T; J Chem Soc, Perkin Trans 1 1999, P1285
(6) Chang, C; Int J Pept Protein Res 1980, V15, P59 HCAPLUS
(7) Inukai, M; Antimicrob Agents and Chemother 1993, V37, P980 HCAPLUS
(8) Isono, F; Antimicrob Agents Chemother 1991, V35, P234 HCAPLUS
(9) Isono, F; J Antibiot 1989, V42, P667 HCAPLUS
(10) Isono, F; J Antibiot 1989, V62, P674
(11) Kaiser, E; Anal Biochem 1970, V34, P595 HCAPLUS
(12) Lee, V; Med Res Rev 1999, V19, P521 HCAPLUS
(13) Merrifield, R; J Org Chem 1993, V58, P5167
(14) Niccolai, D; Chem Commun 1997, P2333 HCAPLUS
(15) Pedroso, E; Tetrahedron Lett 1993, V34, P2195
(16) Raju, B; Bioorg Med Chem Lett 1998, V8, P3043 HCAPLUS
(17) Roth, V: Emerging Therapeutic Targets 1999, V3, P73 HCAPLUS
(18) Schollkopf, U; Angew Chem, Int Ed Engl 1981, V20, P798
(19) Setti, E; Drugs Future 1997, V22, P271 HCAPLUS
(20) Williams, R; Synthesis of Optically Active .alpha.-Amino Acids 1989
IT
     327184-80-5
     RL: RCT (Reactant)
        (prepn. of a combinatorial library of mureidomycin analogs using
                                   solid-phase synthesis)
```

andrick to the contract of the

1.1

RN 327184-80-5 HCAPLUS

CN D-Phenylalanine, N-[(4-nitrophenoxy)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

## IT 327184-69-0P 327184-70-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of a combinatorial library of mureidomycin analogs using solid-phase synthesis)

RN 327184-69-0 HCAPLUS

CN D-Phenylalanine, 3-(1,1-dimethylethoxy)-N-[(4-nitrophenoxy)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA:INDEX:NAME)

Absolute stereochemistry.

RN 327184-70-3 HCAPLUS

CN L-Phenylalanine, 3-(1,1-dimethylethoxy)-N-[(4-nitrophenoxy)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L20 ANSWER 3 OF 12 HCAPLUS COPYRIGHT 2001 ACS

AN 2000:421093 HCAPLUS

DN 133:43809

TI Preparation of new biphenyl and biphenyl-analogous compounds as integrin antagonists

IN Albers, Markus; Urbahns, Klaus; Vaupel, Andrea; Harter, Michael; Schmidt,

```
Delf; Stelte-ludwig, Beatrix; Gerdes, Christoph; Stahl, Elke; Keldenich,
     Jorg; Bruggemeier, Ulf; Lustig, Klemens
     Bayer Aktiengesellschaft, Germany; et al.
PA
     PCT Int. Appl., 360 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
IC
     ICM C07C311-19
     ICS C07C275-42; C07C311-10; C07C311-47; C07D213-40; C07D213-75;
          C07D235-30; A61\mathring{R}031-18; A61\mathring{R}031-4\mathring{4}; A61\mathring{R}031-4184; A61P035-00;
          A61P019-10; A61P027-02
CC
     34-2 (Amino Acids, Peptides, and
     Section cross-reference(s): 1, 25
     PATENT NO.
                       KIND
                              DATE
                                              APPLICATION NO.
PΙ
     WO 2000035864
                        A1
                              20000622
                                              WO 1999-EP9843
                                                                19991213
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
              CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
              IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA,
             MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,
              SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
              CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                            19981216
PRAI US 1998-213381
     MARPAT 133:43809
OS
     Biphenylyl compds. R102CCHR2-U-V-A-B-W-NR3-C-R4 [R1 = H, (un)substituted
AΒ
     alkyl, cycloalkyl, aryl, or (un) satd. heterocyclyl; R2 = H,
     (un) substituted alkyl, cycloalkyl, aryl, or (un) satd. heterocyclyl,
     alkenyl, alkynyl, -NR2'SO2R2'', -NR2'CO2R2', -NR2'COR2', -NR2'CONR2'2, -NR2'CSNR2'2 (R2' has same definition as R1 and R2'' has same definition
     as R1 except it is not H); U or W is a direct bond or (un) substituted
     alkylene; V = (un)substituted alkylene, -NR2'CO- or NR2'SO2-; A and B =
     (un) substituted 1,3- or 1,4-bridging phenylene group or a 2,4- or
     2,5-bridging thienylene group, each of which may have substituents; C is a
     direct bond, CMe(:X-R5)-Y-N(R6)- (R5 is absent, H, (un)substituted alkyl
     or cycloalkyl, NO2, acyl, carboxylic or carboxylate group or is connected
     to R3, Y, R4 or R6, if present; R6 is H, (un) substituted alkyl,
     cycloalkyl, aryl, or {(un)satd. heterocyclyl, an alkylamine or alkylamide
     residue, or is connected to one of R3, R4, Y, or R5, if present, to form a
     heterocyclic ring system; X = CHNO2, CHCN, O, N or S; Y is a direct bond
     or (un) substituted alkylene or alkyne group) or 3,4-dioxo-1,2-
     cyclobutenediyl-NR6-; R3, R4 = H, (un) substituted alkyl, cycloalkyl, aryl,
     or (un)satd. heterocyclyl, an alkylamine or alkylamide residue, or is
     connected to one of R4 (or R3); Yir R53 or 1R6, if present, to form a
     heterocyclic ring system] were prepd. as integrin antagonists.
     (2R,S)-3-[3-(pyridin+3-ylmethylureido)biphenyl-4-yl]-2-[2,4,6-
     trimethylbenzenesulfonylaminolpropanoic acid, prepd. by reactions of
     resin-bound (2R,S)-3-(4-bromophenyl)-2-(9-fluorenylmethoxycarbonylamino)pr
     opanoic acid with sulfonylating, boronic acid, and amine reagents
     (mesitylenesulfonyl chloride, 3-nitrobenzeneboronic acid, and
     2-aminomethylpyridine), showed IC50 = 5 nM for binding to the
     .alpha.v.beta.3 receptor and IC50 = 480 nM in the smooth muscle cell
     migration test.
ST
     amino acid biphenylyl deriv preph antagonist integrin
IT
     Angiogenesis
                                     In the state of
     Antitumor agents
                                     John Carlo
```

Laboratory Fall

一封 郭忠行制设定 [2]

```
Arteriosclerosis
     Eye, disease
     Osteoporosis
     Rheumatoid arthritis
        (prepn. of new biphenyl and biphenyl-analogous compds. as integrin
        antagonists)
ΙT
    Amino acids, preparation
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (prepn. of new biphenyl and biphenyl-analogous compds. as integrin
        antagonists)
    Artery, disease
        (restenosis; prepn. of new biphenyl and biphenyl-analogous compds. as
        integrin antagonists)
IT
     Integrins
     RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (.alpha.v.beta.3; prepn. of new biphenyl and biphenyl-analogous compds.
        as integrin antagonists)
                                    276257-77-3P
                                                    276257-78-4P
                                                                    276257-79-5P
                    276257-76-2P
IT
     276257-75-1P
                    276257-81-9P
                                    276257-82-0P
                                                    276257-83-1P
                                                                    276257-84-2P
     276257-80-8P
                                    276257-87-5P
                    276257<del>,</del>86-4P
                                                    276257-88-6P
                                                                    276257-89-7P
     276257-85-3P
                                                                    276257-94-4P
     276257-90-0P
                    276257-91-1P
                                    276257-92-2P
                                                   276257-93-3P
                    276257-96-6P
                                    276257-97-7P
                                                   276257-98-8P
                                                                    276257-99-9P
     276257-95-5P
                                                                    276258-04-9P
                    276258-01-6P
                                    276258-02-7P
                                                    276258-03-8P
     276258-00-5P
                                                 276258-08-3P
     276258-05-0P
                    276258-06-1P
                                    276258-07-2P
                                                                    276258-09-4P
     276258-10-7P
                    276258-11-8P
                                    276258-12-9P
                                                    276258-13-0P
                                                                    276258-14-1P
                    276258-16-3P
                                    276258-17-4P
                                                    276258-18-5P
                                                                    276258-19-6P
     276258-15-2P
                                    276258-22-1P
                                                    276258-23-2P
                                                                    276258-24-3P
     276258-20-9P
                    276258-21-0P
                                    276258-27-6P 276258-28-7P
     276258-25-4P
                    276258-26-5P
                                                                    276258-29-8P
                                                                    276258-34-5P
     276258-30-1P
                    276258-31-2P
                                    276258-32-3P
                                                    276258-33-4P
                    276258-36-7P
                                    276258-37-8P
                                                    276258-38-9P
                                                                    276258-39-0P
     276258-35-6P
                    276258-41-4P
                                    276258-42-5P
                                                    276258-43-6P
                                                                    276258-44-7P
     276258-40-3P
                    276258-46-9P
                                    276258-47-0P
                                                    276258-48-1P
                                                                    276258-49-2P
     276258-45-8P
                                    276258-52-7P
                                                    27,6258-53-8P
                                                                    276258-54-9P
                    276258#51-6P
     276258-50-5P
                    276258-56-1P
                                                    276258-58-3P
                                                                    276258-59-4P
     276258-55-0P
                                    276258-57-2P
                    276258-61-8P
                                    276258-62-9P
                                                    276258-63-0P
                                                                    276258-64-1P
     276258-60-7P
     276258-65-2P
                    276258-66-3P
                                    276258-67-4P
                                                    276258-68-5P
                                                                    276258-69-6P
     276258-70-9P
                    276258-71-0P
                                    276258-72-1P
                                                    276258-73-2P
                                                                    276258-74-3P
                                                                    276258-82-3P
     276258-75-4P
                    276258-76-5P
                                    276258-79-8P
                                                    276258-80-1P
     276258-83-4P
                    276258-84-5P
                                    276258-85-6P
                                                    276258-88-9P
                                                                    276258-94-7P
                    276259-01-9P
                                    276259-02-0P
                                                    276259-04-2P
                                                                    276259-07-5P
     276258-97-0P
                    276259-13-3P
                                    276259-15-5P
                                                    276259-17-7P
                                                                    276259-22-4P
     276259-10-0P
                                                    276259-29-1P
                                                                    276259-30-4P
     276259-26-8P
                    276259-27-9P
                                    276259-28-0P
                    276259-32-6P
                                    276259-33-7P
                                                    276259-34-8P
                                                                    276259-35-9P
     276259-31-5P
                    276259-37-1P
     276259-36-0P
                                    276259-38-2P
                                                    276259-39-3P
                                                                    276259-40-6P
                                                    276259-44-0P
                                                                    276259-45-1P
     276259-41-7P
                    276259-42-8P
                                    276259-43-9P
                                                    276259-49-5P
                                                                    276259-50-8P
     276259-46-2P
                    276259-47-3P
                                    276259-48-4P
                                                                    276259-55-3P
                                    276259-53-1P
                                                    276259-54-2P
     276259-51-9P
                    276259-52-0P
                                                    276259-59-7P
                                                                    276259-60-0P
     276259-56-4P
                     276259-57-5P
                                    276259-58-6P
                                                                    276259-65-5P
     276259-61-1P
                    276259-62-2P
                                    276259-63-3P
                                                    276259-64-4P
     276259-66-6P
                    276259-67-7P
                                    276259-68-8P
                                                    276259-69-9P
                                                                    276259-70-2P
                     276259-72-4P
                                    276259-73-5P
                                                    276259-74-6P
                                                                    276259-75-7P
     276259-71-3P
     276259-76-8P
                    276259-77-9P
                                    276259-78-0P
                                                    276259-79-1P
                                                                    276259-80-4P
                                    276259-83-7P
                                                    276259-84-8P
                                                                    276259-85-9P
     276259-81-5P
                    276259-82-6P
                                                                    276259-90-6P
     276259-86-0P
                     276259-87-1P
                                    276259-88-2P
                                                    276259-89-3P
     276259-91-7P
                    276259-92-8P
                                    276259-93-9P
                                                    276259-94-0P
                                                                    276259-95-1P
                    276259-97-3P
                                    276259-98-4P
                                                    276259-99-5P
                                                                    276260-00-5P
     276259-96-2P
                                                                    276260-05-0P
                    276260-02-7P
                                    276260-03-8P
                                                    276260-04-9P
     276260-01-6P
                                                                    276260-10-7P
                    276260-07-2P
                                    276260-08-3P
                                                    276260-09-4P
     276260-06-1P
```

```
276260-12-9P
                                    276260-13-0P
                                                    276260-14-1P
                                                                   276260-15-2P
    276260-11-8P
                                                                   276260-20-9P
                    276260-17-4P
                                    276260-18-5P
                                                    276260-19-6P
    276260-16-3P
                                                    276260-24-3P
                                                                   276260-25-4P
    276260-21-0P
                    276260-22-1P
                                    276260-23-2P
                                  276260-28-7P 276260-29-8P
    276260-26-5P 276260-27-6P
                                    276260-32-3P 276260-33-4P
                                                                   276260-34-5P
                    276260-31-2P
    276260-30-1P
                                                                   276260-39-0P
                    276260 536-7P
                                    276260-37-8P
                                                    276260-38-9P
    276260-35-6P
    276260-40-3P
    RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
    preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (prepn. of new biphenyl and biphenyl-analogous compds. as integrin
        antagonists)
                    276260-42-5P
                                    276260-43-6P 276260-44-7P
                                                                   276260-45-8P
IT
     276260-41-4P
                                   276260-48-1P
                    276260-47-0P
                                                    276260-49-2P
                                                                   276260-50-5P
     276260-46-9P
                                    276260-53-8P
                                                    276260-54-9P
                                                                   276260-55-0P
                    276260-52-7P
     276260-51-6P
                                                    276260-59-4P
                                                                   276260-60-7P
                    276260-57-2P
                                    276260-58-3P
     276260-56-1P
                                                   276260-64-1P
                    276260-62-9P
                                    276260-63-0P
                                                                   276260-65-2P
     276260-61-8P
                                                                   276260-70-9P
     276260-66-3P
                    276260-67-4P
                                    276260-68-5P
                                                    276260-69-6P
                                                                   276260-75-4P
                                    276260-73-2P
                                                    276260-74-3P
     276260-71-0P
                    276260-72-1P
                                                    276260-79-8P
                                                                   276260-80-1P
                    276260-77-6P
                                    276260-78-7P
     276260-76-5P
                                    276260-83-4P
                                                    276260-84-5P
                                                                   276260-85-6P
     276260-81-2P
                    276260-82-3P
     276260-86-7P
                    276260-87-8P
                                    276260-88-9P
                                                    276260-89-0P
                                                                   276260-91-4P
     276260-93-6P
                    276260-95-8P
                                    276260-96-9P
                                                    276260-97-0P
                                                                    276260-98-1P
                                    276261-01-9P
                                                    276261-02-0P
                                                                    276261-03-1P
     276260-99-2P
                    276261-00-8P
     276261-04-2P
                    1276261÷05-3P
                                    276261-06-4P
                                                    276261-07-5P
                                                                    276261-08-6P
                                                                    276261-15-5P
     276261-09-7P
                    276261-10-0P
                                    276261-12-2P
                                                    276261-14-4P
                    276261-19-9P
                                    276261-21-3P
                                                    276261-22-4P
                                                                    276261-23-5P
     276261-17-7P
                    276261-25-7P
                                                    276261-27-9P
                                                                    276261-28-0P
     276261-24-6P
                                    276261-26-8P
                    276261-30-4P
                                    276261-31-5P
                                                    276261-32-6P
                                                                    276261-33-7P
     276261-29-1P
                                                                    276261-38-2P
                    276261-35-9P
                                    276261 - 36 - 0P + 276261 - 37 - 1P
     276261-34-8P
                                    276261-43-9P
                                                    276261+45-1P
                                                                   276261-47-3P
                    276261-41-7P
     276261-39-3P
                    276261-50-8P
                                    276261-51-9P 276261-53-1P
                                                                   276261-54-2P
     276261-48-4P
     276261-56-4P
                    276261-57-5P
                                    276261-58-6P
                                                    276261-60-0P
                                                                    276261-62-2P
                                    276261-67-7P
                                                    276261-69-9P
                                                                    276261-71-3P
     276261-64-4P
                    276261-65-5P
                                    276261-75-7P
                                                    276261-76-8P
                                                                    276261-77-9P
                    276261-74-6P
     276261-72-4P
                    276261-79-1P
                                    276261-80-4P
                                                    276261-81-5P
                                                                    276261-82-6P
     276261-78-0P
     276261-83-7P
                                    276261-85-9P
                                                    276261-86-0P
                                                                    276261-87-1P
                    276261-84-8P
                                    276261-90-6P
                                                    276261-91-7P
                                                                    276261-92-8P
     276261-88-2P
                    276261-89-3P
     276261-93-9P
                    276261-94-0P
                                    276261-95-1P
                                                    276261-96-2P
                                                                    276261-97-3P
                                                    276262-01-2P
                                                                    276262-02-3P
     276261-98-4P
                    276261-99-5P
                                    276262-00-1P
                                    276262-05-6P
                                                    276262-06-7P
                                                                    276262-07-8P
     276262-03-4P
                    276262-04-5P
                    276262-09-0P
                                    276262-10-3P
                                                    276262-11-4P
                                                                    276262-12-5P
     276262-08-9P
                    276262-14-7P
                                    276262-15-8P
                                                    276262-16-9P
                                                                    276262-17-0P
     276262-13-6P
                                                    276262-21-6P
                                                                    276262-22-7P
                    276262-19-2P
                                    276262-20-5P
     276262-18-1P
                                    276262-25-0P
                                                                    276262-27-2P
     276262-23-8P
                    276262-24-9P
                                                    276262-26-1P
                    276262-29-4P
                                    276262-30-7P
                                                    276262-31-8P
                                                                    276262-32-9P
     276262-28-3P
                    276262-34-1P
                                    276262-35-2P
     276262-33-0P
                                                    276262-36-3P
                                                                    276262-37-4P
                                    276262-40-9P
                                                    276262-41-0P
                                                                    276262-42-1P
     276262-38-5P
                    276262-39-6P
     276262-43-2P
                    276262-44-3P
                                    276262-45-4P
                                                    276262-46-5P
                                                                    276262-47-6P
                                    276262-50-1P
                                                    276262-51-2P
                                                                    276262-52-3P
     276262-48-7P
                    276262-49-8P
                                    276262-55-6P
                                                    276262-56-7P
                                                                    276262-57-8P
                    276262-54-5P
     276262-53-4P
                                                                    276262-62-5P
     276262-58-9P
                                    276262-60-3P
                                                    276262-61-4P
                    276262-59-0P
                                                                    276262-67-0P
                                    276262-65-8P
                                                    276262-66-9P
     276262-63-6P
                    276262-64-7P
                                        The state of
     276262-68-1P
                    276262-69-2P
     RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
     (Preparation); USES (Uses)
        (prepn. of new biphenyl and biphenyl-analogous compds. as integrin
        antagonists)
     51-45-6, 2-(Imidazol-4-yl)ethylamine, reactions
                                                         78-81-9, Isobutylamine
IT
```

; ;-

```
88-14-2, 2-Furancarboxylic acid 96-15-1, 2-Methylbutylamine
                        98-58-8, 4-Bromobenzenesulfonyl chloride
                                                                     98-60-2,
     2-Aminothiazole
     4-Chlorobenzenesulfonyl chloride 107-15-3, 1,2-Ethanediamine, reactions 108-00-9, n,n-Dimethylethylenediamine 108-91-8, Cyclohexylamine, reactions 462-08-8, 3-Aminopyridine 504-24-5, 4-Aminopyridine
     504-29-0, 2-Aminopyridine 645-36-3, Aminoacetaldehyde diethyl acetal
     765-30-0, Cyclopropylamine
                                  773-64-8, 2,4,6-Trimethylbenzenesulfonyl
                 934-32-7, 2-Aminobenzimi dazole
                                                    1001-53-2,
     n-Acetylethylenediamine 1003-03-8, Cyclopentylamine
                                                                 2905-23-9,
     2-Chlorobenzenesulfonyl chloride 2905-24-0, 3-Bromobenzenesulfonyl
                2991-42-6, 4-Trifluoromethylbenzenesulfonyl chloride
     3731-51-9, 2-Aminomethylpyridine 3731-52-0, 3-Aminomethylpyridine
     3731-53-1, 4-Aminomethylpyridine 4548-45-2, 2-Chloro-5-nitropyridine
     4659-45-4, 2,6-Dichlorobenzoyl chloride 4795-29-3, 2-
     Aminomethyltetrahydrofuran
                                    5231-87-8; 5402-73-3, 2,5-
     Dichlorobenzenesulfonyl chloride 6335-76-8
                                                       7154-73-6,
     2-Pyrrolidin-1-ylethylamine
                                     7693-46-1, 4-Nitrophenyl chloroformate
     10191-60-3, Dimethyl cyanimidodithiocarbonate: 13258-63-4,
                            13331-27-6 13623-94-4 13952-84-6,
     4-Pyridineethanamine
                       20781-20-8, 2,4 Dimethoxybenzylamine
                                                                 21286-54-4, +
     sec-Butylamine
     Camphor 10 sulfonyl chloride 22374-89-6 23095-05-8, 5-Bromo-2-methoxybenzenesulfonyl chloride 29022-11-5, Fmoc gly oh
                   66472-86-4 79711-73-2 79844-65-8
                                                           80500-27-2
     50998-05-5
     87199-16-4, 3-Formylbenzeneboronic acid 87199-17-5, 4-
                                 88831-43-0
276262-70-5 | 276262-71-6
     Formylbenzeneboronic acid
                                                               107819-90-9
     126727-04-6
                    180181-93-5
                                     RL: RCT (Reactant)
         (prepn. of new biphenyl and biphenyl-analogous compds. as integrin
                                     Harris de 1
        antagonists)
IT
     276258-77-6P
                     276258=78-7P
                                    276258-81-2P 276258-86-7P
                                                                     276258-87-8P
                                     276258-91-4P 276258-92-5P
     276258-89-0P
                    276258-90-3P
                                                                     276258-93-6P
                     276258-96-9P
                                    276258-98-1P
                                                     276258-99-2P
                                                                     276259-00-8P
     276258-95-8P
                                     276259-06-4P 276259-08-6P
                                                                     276259-09-7P
                     276259-05-3P
     276259-03-1P
                                                                     276259-18-8P
                     276259-12-2P
                                     276259-14-4P
                                                     276259-16-6P
     276259-11-1P
                                    276259-21-3P 276259-23-5P
                                                                     276259-24-6P
                     276259-20-2P
     276259-19-9P
                                     276262-73-8P
     276259-25-7P
                     276262-72-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. of new biphenyl and biphenyl-analogous compds. as integrin
                                    明國於蘇爾市
        antagonists) 🕟
                                     RE.CNT
(1) Anon; WO 9736859 A 1997 HCAPLUS
(2) Merck, P; DE 19548709 A 1997 HCAPLUS 18 18
(3) Merck, P; WO 9800395 A 1998 HCAPLUS
(4) Tanabe, S; WO 9936393 A 1999 HCAPLUS
                                    一种 建甲基甲基苯甲基苯
IT
     276260-27-6P
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
     276260-27-6P
        (prepn. of new biphenyl and biphenyl-analogous compds. as integrin
                                        antagonists)
     276260-27-6 HCAPLUS
RN
     [1,1'-Biphenyl]-4-propanoic acid, alpha.-[[(4-
CN
     methoxyphenoxy) carbonyl]amino]-3"-[[(propylamino)carbonyl]amino]-,
     (.alpha.S) - (9CI) (CA INDEX NAME)
```

国集品的4°53

```
n-PrNH H CO2H O OMe
```

ANSWER 4 OF 12 HCAPLUS COPYRIGHT 2001 ACS

```
AN
     1999:584481 HCAPLUS
DN
     132:12483
     Enantiomeric Separation of (.+-.) Amino Acid Esters on
ΤI
     (-)-Phenylurea Chiral Stationary, Phase
     Lee, Kwang-Pill; Lee, Hyun-Bong; Lee, Young Cheol; Choi, Seong-Ho; Ryoo,
ΑU
     Jae Jeong; Park, Jung Hag
     Department of Chemistry, Graduate School, Kyungpook National University,
CS
     Sangeok-dong, Taegu, 702-701, S. Korea
     Microchem. J. (1999), 63(1), 18-23
SO
     CODEN: MICJAN; ISSN: 0026-265X
PB
     Academic Press
DT
     Journal
LΑ
     English
CC
     34-2 (Amino Acids, Peptides, and
     Proteins)
     Section cross-reference(s): 9 to let
AΒ
     The 3,5-dinitrobenzoyl (.+-.)-amino acid esters were successfully resolved
     on (-)-phenylurea chiral stationary phases (CSPs) in a normal phase mode
     by high-performance liq. chromatog. (HPLC). The alcs. used for
     esterification were methanol, ethanol, and n-propanol. The effects of
     esterification were studied via retention and optical resoln. The solvent
     and its concn. effect on enantioselectivity have been
     investigated based on the binary or ternary solvent system. The alc. used
     in the binary or ternary solvent system was crit. to the
     enantiomeric resoln. of 3,5-dinitrobenzoyl amino acid esters while the nonalcoholic solvent was not suitable. The optical condition of the
     enantiomeric resoln. is discussed in terms of the solvent compn.
     and structure of the amino acid esters The main chiral recognition
     mechanism based on the .pi.-.pi. interaction of the nitrobenzoyl group of
     the amino acid derivs. with the 'pi.-basic Ph group of CSPs is described.
     (c) 1999 Academic Press.
ST
     amino acid ester enantiomeric sepn phenylurea chiral
     stationary phase; chromatog chiral stationary phase sepn amino
                                    Figure 4 标
     acid ester enantiomeric
IT
     Chiral recognition : # | - |
     Chromatography
     Esterification
                                   · 5.25
     Resolution (separation)
        (enantiomeric sepn; of (.+-:) amino acid esters on
        (-)phenylurea chiral stationary phase)
IT
     Amino acids, preparation
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (enantiomeric sepn. of (.++:) amino acid esters on
        (-)phenylurea chiral stationary phase) 🐔
                                  IT
     64-10-8, Phenylurea
     RL: RCT (Reactant)
        (chiral stationary phase for the resoln: of (.+-.)-amino acid esters)
                                   医骶骨髓 医胚胎
```

THE PROPERTY OF

L20

```
74928-20-4 74928-21-5 74928-22-6 104336-95-0 106145-09-9 135088-7
     35519-07-4
                     74928-18-0
IT
                     92915-46-3
                                                                      135088-75-4
     74928-24-8
                                       135449-39-7 138088-19-4
     135088-79-8
                     135449-37-5
                                                                      138088-20-7
                      138088-22-9
                                       148346-54-7
                                                        187836-09-5
                                                                         214782-91-9
     138088-21-8
                                       25153845048 - 251538-51-9
                      251538-49-5
                                                                         251538-52-0
      251538-48-4
     251538-54-2
                      251538+55-3
                                       251538-56-4
                                                       251538-57-5
                                                                         251538-58-6
     251538-59-7 251538-60-0 251538-61-1 251538-62-2 251538-64-4 251538-65-5 251538-66-6 251538-67-7 251538-68-8 251538-69-9
                                                                         251538-63-3
                                       251538-70-2 251538-71-3
                      251538-69-9
     251538-68-8
                                         251538-73-5
     251538-72-4
     RL: RCT (Reactant)
                              1 .
         (enantiomeric sepn. of on (-) phenylurea chiral
         stationary phase)
                                         - I. O. O. - 10
RE.CNT
RE
(1) Armstrong, D; J Chromatogr 1991, V539, P83 HCAPLUS
(2) Gisch, D; The 9th International Symposium on Column Liquid Chromatography
    1985
(3) Keith, J; Chirality 1993, V5, P201
(4) Lam, S; J Chromatogr Sci 1984, V22, P416 HCAPLUS
(5) Okamoto, Y; Bull Chem Soc Jpn 1990, V63, P955 HCAPLUS
(6) Okamoto, Y; Chem Lett 1987, P1857 HCAPLUS (7) Okamoto, Y; Chemtech 1988, P176 HCAPLUS (8) Okamoto, Y; J Chromatogr 1986, V363, P173 HCAPLUS
(9) Pirkle, W; J Am Chem Soc 1987, V109, P5975 HCAPLUS (10) Pirkle, W; J Chromatogr 1984, V316, P585 HCAPLUS
(11) Pirkle, W; J Org Chem 1992, V57, P3854 HCAPLUS (12) Ryu, J; Anal Sci 1997, V13, P217 HCAPLUS
(13) Ryu, J; J Chromatogr A 1998, V14, 2247

IT 251538-65-5 251538-68-8 251538-71-3
     RL: RCT (Reactant)
         (enantiomeric sepn. of on (-) phenylurea chiral
         stationary phase)
RN
     251538-65-5 HCAPLUS
     Valine, N-[(1-naphthalenyloxy)carbonyl]-, methyl ester (9CI) (CA INDEX
CN
MeO-C
i-Pr-CH-NH-C-O
     251538-68-8 HCAPLUS
RN
     Phenylalanine,: N-[(1-naphthalenyloxy)carbonyl]-, methyl ester (9CI) (CA
CN
```

April 1 1

INDEX NAME)

RN 251538-71-3 HCAPLUS

CN Tryptophan, N-[(1-naphthalenyloxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

offic h.

L20 ANSWER 5 OF 12 HCAPLUS COPYRIGHT 2001 ACS

AN 1999:35812 HCAPLUS

DN 130:153966

TI Solid-phase synthesis of tyrosine peptide aldehydes. Analogs of (S)-MAPI

AU Page, Patrick; Bradley, Mark; Walters, Iain; Teague, Simon

CS Department of Chemistry, University of Southampton, Southampton, SO17 1BJ, UK

SO J. Org. Chem. (1999), 64(3), 794+799 CODEN: JOCEAH; ISSN: 0022-3263

PB American Chemical Society

DT Journal

LA English

CC 34-3 (Amino Acids, Peptides, and Proteins)

We report an efficient solid-phase synthesis of C-terminal tyrosine AB peptide aldehydes based on the HIV protease inhibitors (S)-MAPI and GE 20372 A. Our strategy consisted of anchoring the side chain of Dde-Tyrosinol onto the brominated Wang linker deriv. ((4-bromomethyl)phenoxy-allyl acetate) to give after ester hydrolysis the N.alpha.-(Dde)-O-(4-methylphenoxyacetic acid)-L-Tyrosinol template. was attached to aminomethyl resin and elongated using std. Fmoc protocols. Importantly there was no evidence of esterification side reactions. The unsym. substituted urea linkage of the (S)-MAPI family was incorporated using the N.alpha.-(4-nitrophenyloxycarbonyl)amino acid tert-Bu esters following which the protected tetrapeptide alc. immobilized on the solid support was oxidized to its corresponding aldehyde using sulfur trioxide-pyridine. The efficiency and reliability of the oxidn. step was dramatically improved by the incorporation of a small PEG-spacer between the linker and the solid support. The tetrapeptides were cleaved by acidolysis, purified by RP HPLC, and isolated in high yield and purity, demonstrating the success of the whole synthetic process.

ST solid phase synthesis tyrosine peptide aldehyde oxidn

IT Peptides, preparation!

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic

.a 'd

Timothy Saunders EIC-LAW, Lib. 308-413

i de glidere prije i ka Nadrije je je ja prije Nagreje i koji ki

THE THE A ST. P. STAFF

```
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
                                   . 1. 1
                                          (Preparation); USES (Uses)
         (aldehydes; solid-phase synthesis of tyrosine peptide aldehydes)
                                      विकास हो। स
IT
     Oxidation
                                      644
     Solid phase synthesis
         (solid-phase synthesis of tyrosine peptide aldehydes)
IT
     70857-49-7P
                    163565-75-1P, GE 20372a
     RL: BAC (Biological activity or effector, except adverse); PNU
     (Preparation, unclassified); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
         (solid-phase synthesis of tyrosine peptide aldehydes)
                     220237-28-5P
IT
     220237-27-4P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
         (solid-phase synthesis of tyrosine peptide aldehydes)
IT
     14221-01-3
     RL: CAT (Catalyst use); USES (Uses)
         (solid-phase synthesis of tyrosine peptide aldehydes)
     693-13-0, Diisopropyl carbodiimide 7087-68-5, Diisopropyl ethylamine
IT
     7446-11-9, Sulfur trioxide, reactions 13887-98-4
                                                              57260-73-8
     155505-56-9
                    187526-99-4 191425-55-5 191426-93-4
     RL: RCT (Reactant)
         (solid-phase synthesis of tyrosine peptide aldehydes)
IT
     220237-29-6DP, resin-bound
                                    220237-30-9P
                                                    220237-31-0P
                                                                     220237-32-1P
     220237-33-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (solid-phase synthesis of tyrosine peptide aldehydes)
                                              RE.CNT
                                    · 1411年的新生生
RE
(1) Aoyagi, T; J Antibiot 1969, V22, P283 HCAPLUS
(2) Aoyagi, T; Proteases and Biological Control 1975, P429 HCAPLUS
(3) Bendal, M; Eur J Biochem 1977, V79, P201
(4) Bloomberg, G; Tetrahedron Lett 1993; V34, P4709 HCAPLUS
(5) Bullock, T; J Mol Biol 1996, V255, P714 HCAPLUS
(6) Bycroft, B; J Chem Soc Chem Commun 1993, P778 HCAPLUS
(7) Chen, C; J Am Chem Soc 1994, V116, P2661 HCAPLUS
(8) Dinh, T; Tetrahedron Lett 1996, V37, P1161 HCAPLUS
(9) Ede, N; Tetrahedron Lett 1997, V38, P7119 HCAPLUS
(10) Fehrentz, J; Tetrahedron Lett 1995, V36, P7871 HCAPLUS
(11) Frankfater, A; Biochemistry 1981, V20, P5517 HCAPLUS
(12) Galeotti, N; Lett Pept Sci 1997, V4, P437 HCAPLUS
(13) Kaneto, R; J Antibiot 1993, V46, P1622 HCAPLUS
(14) Kennedy, W; Biochemistry 1979, V18, P349 HCAPLUS (15) Kunz, H; Angew Chem Int Ed Engl. 1984, V23, P436 (16) Mackenzie, N; Biochemistry 1986, V25, P2293 HCAPLUS
(17) Malcolm, B; Biochemistry 1995, V34, P8172 HCAPLUS
(18) Marsh, I; J Org Chem 1997, V62, P6199 HCAPLUS
(19) Murphy, A; J Am Chem Soc 1992, V114, P3156 HCAPLUS (20) Nahm, S; Tetrahedron Lett 1981, V22, P3815 HCAPLUS
(21) Ngu, K; Tetrahedron Lett 1997, V38, P973 HCAPLUS
(22) Ogilvie, W; J Med Chem 1997, V40, P4113 HCAPLUS
(23) Rano, T; Tetrahedron Lett 1995, V36, P3789 HCAPLUS
(24) Richter, L; Tetrahedron Lett 1994, V35, P4705 HCAPLUS
(25) Rotella, D; J Am Chem Soc 1996; V118; P12246 HCAPLUS
(26) Sarin, V; Anal Biochem 1981, V20, P147
(27) Sarubbi, E; FEBS Lett 1993, V319, P253 HCAPLUS
(28) Schacht, A; Bioorg Med Chem Lett 1995, V5, P2529 HCAPLUS
(29) Schroder, E; FEBS Lett 1993, V315, P38 HCAPLUS
(30) Schultz, R; J Biol Chem 1989, V264, P1497 HCAPLUS
```

24 BHT 6

jt: Ni⊩

```
(31) Stefanelli, S; J Antibiot 1995, V48, P332 HCAPLUS
```

- (32) Stella, S; J Antibiot 1991, V44, P1019 HCAPLUS (33) Thompson, R; Biochemistry 1973, V12, P47 HCAPLUS
- (34) Watanbe, T; Agric Biol Chem 1979, V43, P243
- (35) Weinshenker, N; Tetrahedron Lett 1972, V32, P3281
- (36) Westerik, J; J Biol Chem 1972, V247, P8195 HCAPLUS
- (37) Woo, J; Bioorg Med Chem Lett 1995, V5, P1501 HCAPLUS
- (38) Zhang, X; J Org Chem 1997, V62, P6420 HCAPLUS
- ΙT 191425-55-5 191426-93-4

RL: RCT (Reactant)

(solid-phase synthesis of tyrosine peptide aldehydes)

RN191425-55-5 HCAPLUS

L-Phenylalanine, N-[(4-nitrophenoxy)carbonyl]-, 1,1-dimethylethyl ester CN (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191426-93-4 HCAPLUS

L-Tyrosine, O-(1,1-dimethylethyl)-N-[(4-nitrophenoxy)carbonyl]-, CN (CA INDEX NAME) 1,1-dimethylethyl ester (9CI)

Absolute stereochemistry

- ANSWER 6 OF 12 HCAPLUS L20 COPYRIGHT
- 1997:598126 HCAPLUS AN
- 127:176703 DN
- TI Solid-Phase Total Synthesis of Oscillamide Y and Analogs
- Marsh, Ian R.; Bradley, Mark; Teague, Simon J. ΑU
- Department of Chemistry, University of Southampton, Southampton, SO17 1BJ, CS UK
- J. Org. Chem. (1997), 62(18), 6199-6203 SO CODEN: JOCEAH; ISSN: 0022-3263
- American Chemical Society PΒ
- DT Journal
- English LА
- CC 34-3 (Amino Acids, Peptides, Proteins)

GΙ

An efficient solid phase synthesis of oscillamide Y (I) and three analogs AΒ is reported. The cyclic peptide was prepd. using a combination of 9-fluorenylmethoxycarbonyl (Fmoc) and allyl chemistries and an acid labile Wang type linker. The urea functionality was smoothly incorporated using D-lysine building block 4-O2NC6H4O2C-D-Lys(Fmoc)-OCH2CH:CH2. Coupling to the N-Me amino acid was readily achieved using HATU, monitoring the reaction using bromophenol blue. Allyl deprotection was accomplished using Pd(PPh3)4 and dimedone, and cyclization was smoothly accomplished using PyBroP. All reactions were monitored using mass spectrometry methodol. The cyclized materials were cleaved by acidolysis and purified by RP HPLC. In all cases the material isolated was the major product and gave the expected mol, ion information. HPLC comparison with an authentic sample of oscillamide Y showed that the isomer contg. N-methyl-L-alanine and L-homotyrosine was the natural product. 1H NMR and 1H-1H COSY NMR expts. further confirmed this identification. The four compds. were tested as competitive and slow-tight binding inhibitors of chymotrypsin but showed, contrary to literature expectations, no inhibitory activity. ST oscillamide Y isomer solid phase synthesis IT Solid phase peptide synthesis (solid-phase total synthesis of oscillamide Y and analogs) 68858-21-9, 4-(Hydroxymethyl)phenoxyacetic acid IT to be the first of RL: RCT (Reactant) (handle; solid-phase total synthesis of oscillamide Y and analogs) 34404-32-5, D-Lysine, N6-[(phenylmethoxy)carbonyl]-IT RL: RCT (Reactant) (solid-phase total synthesis of oscillamide Y and analogs) 115186-31-7P, Boc-D-Lys (Fmoc)-OH IT 55878-47-2P 193948-47-9P 193948-49-1P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (solid-phase total synthesis of oscillamide Y and analogs) 194038-14-7P 194038-15-8P IT 168482-80-2P, Oscillamide Y RL: SPN (Synthetic preparation); PREP (Preparation) (solid-phase total synthesis of oscillamide Y and analogs) IT 193948-49-1P RL: RCT (Reactant); SPN (Synthétic préparation); PREP (Preparation) (solid-phase total synthesis of oscillamide Y and analogs) 4、野山鄉 博門 RN193948-49-1 HCAPLUS: 4 L-Lysine, N6-[(9H-fluoren-9-ylmethoxy)carbonyl]-N2-[(4-CN nitrophenoxy)carbonyl]-, 2-propenyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

1 10 23 LIEL MINAM GOL

```
2001 ACS
L20
    ANSWER 7 OF 12
                    HCAPLUS
                              COPYRIGHT
AN
     1996:610346 HCAPLUS
DN
     125:301598
    A solid pharmaceutical composition providing improved oral bioavailability
TI
     for HIV protease inhibitors
IN
    Al-razzak, Laman A.; Marsh, Kennan C.; Pyter, Richard A.
PA
    Abbott Laboratories, USA
     U.S., 18 pp. Cont.-in-part of U.S. Ser. No. 267,273, abandoned.
SO
     CODEN: USXXAM
DT
     Patent
LA
     English
IC
     ICM A61K031-16
        A61K031-695; A61K031-425; A61K031-42
NCL
     514616000
     34-2 (Amino Acids, Peptides,
     Proteins)
     Section cross-reference(s): 63 page 1983
FAN.CNT 2
                      KIND:
                            DATE
                                           APPLICATION NO.
                           19950413 WO 1994-WO101
PΙ
    US 5559158
                            19960924
                                          US 1994-297004
                                                            19940831
    CA 2167413
                      AA :
                                                            19940909
    WO 9509614
                      A1 ,
                                                            19940909
            AU, CA, JP, KR
        RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
    AU 9477229
                       A1 19950501 AU 1994-77229
                                                            19940909
                      B2 19980122 A WAR A A
    AU 685509
                         19960717 EP 1994-928043
20010328 EP 1994-928043
    EP 721330
                     : A1
                                                            19940909
                      B1 20010328
     EP 721330
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
     JP 09503501
                      T2 19970408
                                         JP 1994-510810
                                                            19940909
                            20010415
                                           AT 1994-928043
                                                            19940909
    AT 200023
                       Ε
    IL 110915
                                           IL 1994-110915
                                                            19940911
                            19991222
                      A1
                      A 19970311
                                          US 1996-650261
    US 5610193
                                                            19960522
PRAI US 1993-130409
                            19931001:
                      B2 ·
                       B2 | 19940628
    US 1994-267273
    US 1994-297004
                            19940831
                      W . .
    WO 1994-US10096
                            19940909
    US 1995-424740
                       В1
OS
    MARPAT 125:301598
```

开制量 : 4 (中)

美数数性与标准主义 禁忌

jarinsydder i by Tropins (b. 187

A solid pharmaceutical compn. is claimed, comprising a pharmaceutically AΒ acceptable adsorbent or a mixt. of pharmaceutically acceptable adsorbents to which is adsorbed a mixt. of (1) a pharmaceutically acceptable org. solvent or a mixt. of pharmaceutically acceptable org. solvents, (2) HIV protease inhibitor I, and (3) a pharmaceutically acceptable acid or a combination of pharmaceutically acceptable acids. Thus, e.g., a capsule compn. contg. (% by wt.): (all-S)-I = (2S,3S,5S)-5-[N-[N-[[N-methyl-N-[(2-methyl-N-[(3-methyl-N-[(2-methylisopropyl-4-thiazolyl)methyl]amino]carbonyl]valinyl]amino]-2-[N-[(5thiazolyl)methoxycarbonyl]amino]-1,6-diphenyl-3-hydroxyhexane (prepn. given) (21.84); propylene glycol, USP (10.96); Ethanol, dehydrated USP, 200 proof (22.99); polysorbate 80, NF (5.31); Cremophor EL (4.4); HCl, reagent grade (1.18); Cab-o-sil (26.88) exhibited 89.6 mean % oral bioavailability in dogs vs. <2.0 for unformulated (all-S)-I in capsules. ST oral pharmaceutical dosage HIV protease inhibitor

Ι

الإستراطية

Acquired immune deficiency syndrome IT

(solid pharmaceutical compn. providing improved oral bioavailability for HIV protease inhibitors)

ΙT Castor oil

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (ethoxylated, component of pharmaceutical formulation; solid pharmaceutical compn. providing improved oral bioavailability for HIV the one district of protease inhibitors)

Pharmaceutical dosage forms IT

> (oral, solid pharmaceutical compn. providing improved oral bioavailability for HIV protease inhibitors)

50-81-7, Ascorbic acid, biological studies 57-55-6, Propylene glycol, IT 64-17-5, Ethanol, biological studies 104-15-4, biological studies p-Toluenesulfonic acid, biological studies 7631-86-9, Silicon dioxide, biological studies 7647-01-0; Hydrochloric acid, biological studies 14807-96-6, Talc, biological studies 9005-65-6, Polysorbate 80 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(component of pharmaceutical formulation; solid pharmaceutical compn. providing improved oral bioavailability for HIV protease inhibitors)

9004-34-6, Cellulose, biological studies 1

d)

RL: THU (Therapeutic use); BIOL (Biological; study); USES (Uses) (microcryst., component of pharmaceutical formulation; solid pharmaceutical compn. providing improved oral bioavailability for HIV protease inhibitors)

IT 162990-01-4P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic, use); BIOL (Biological study); PREP india y inggi sa (Preparation); USES (Uses)

(solid pharmaceutical compn. providing improved oral bioavailability for HIV protease inhibitors) 自由于中国的 10 mm 19 1

ΙT 144114-21-6, Retropepsin

RL: BSU (Biological study, unclassified); MSC (Miscellaneous); BIOL

·产工的规则, 为工作。 . The standar

प्रकार है। इस्मान ने इस् 

IT

```
district.
                                     (Biological study)
         (solid pharmaceutical compn. providing improved oral bioavailability
         for HIV protease inhibitors)
IT
                     144164-10-3P
     143838-10-2P
     RL: BYP (Byproduct); PREP (Preparation)
         (solid pharmaceutical compn. providing improved oral bioavailability
         for HIV protease inhibitors)
     75-12-7, Formamide, reactions 105-39-5, Ethyl chloroacetate 534-07-6, 1,3-Dichloroacetone 563-83-7, Isobutyramide 6306-52-1, L-Valine methyl
IT
     ester hydrochloride 6372-14-17 N= (Benzylokycarbonyl)-L-phenylalaninol
     7693-46-1, 4-Nitrophenyl chloroformate 124424-99-5, Di-tert-butyl
     dicarbonate 153441-77-1 156732-13-7 156732-15-9
     RL: RCT (Reactant)
         (solid pharmaceutical compn. providing improved oral bioavailability
         for HIV protease inhibitors)
ΙT
     115-08-2P, Thioformamide
                                  13515-65-6P, Thioisobutyramide
                                                                     32955-21-8P,
     2-Amino-5-(ethoxycarbonyl)thiazole 32955-22-9P, Ethyl thiazole-5-carboxylate 33142-21-1P, Ethyl 2-chloro-2-formylacetate
     38585-74-9P, 5-(Hydroxymethyl)thiazole 59830-60-3P, N-(Benzyloxycarbonyl)-L-phenylalaninal 65386-28-9P, 4-(Chloromethyl)-2-
     isopropylthiazole hydrochloride 137649-69-5P
                                                          144141-68-4P
                                     144163-85-9P" 144163-97-3P
     144163-43-9P
                     144163-44-0P
                                                                     144164-11-4P
     154212-59-6P
                     154212-60-9P
                                   154212-61-0P # 154248-99-4P
     162537-10-2P, N-(4-Nitrophenoxycarbonyl)-L-valine Methyl Ester
                                     162849-94-7P
     162849-92-5P
                     162849-93-6P
                                                     162849-95-8P
                                                                     162849-96-9P
     162990-03-6P
                     165315-39-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (solid pharmaceutical compn. providing improved oral bioavailability
        for HIV protease inhibitors)
ΙT
     155213-67-5P
     RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
         (solid pharmaceutical compner providing improved oral bioavailability
        for HIV protease inhibitors)
ΙT
     153441-77-1
     RL: RCT (Reactant)
         (solid pharmaceutical compn. providing improved oral bioavailability
        for HIV protease inhibitors)
RN
     153441-77-1 HCAPLUS
CN
     L-Valine, N-(phenoxycarbonyl)-, methyl ester (9CI)
                                                             (CA INDEX NAME)
Absolute stereochemistry.
                                    Mitthen, at
                           i bing e
                                    ring grade
                           4...
            OPh
    HN
                                    OMe
                           4
```

表介的基本的的特殊。 表示的一个可能的一个。 21章:"想题的表示真实。"

### Absolute stereochemistry.

```
ANSWER 8 OF 12 HCAPLUS COPYRIGHT 2001 ACS
L20
     1995:928117 HCAPLUS
ΑN
     123:340946
DN
ΤI
     Novel amino acid ester and reagent composition for detecting
     leukocytes or elastase in bodily fluids.
                                    2 to 1 to
IN
     Yagi, Yuji
PΑ
     Kyoto Daiichi Kagaku Co., Ltd., Japan
SO
     Eur. Pat. Appl., 19 pp.
     CODEN: EPXXDW
DT
     Patent
     English
LΑ
IC
     ICM C07D295-12
     ICS C07D213-81; C07D213-82; C07D307-68; C07D209-36; C07D213-64;
          C07D311-46; C07C311-19; C07D309-32; C12Q001-44
     34-2 (Amino Acids, Peptides, and
CC
     Section cross-reference(s): 7,
FAN.CNT 1
                                                            DATE
     PATENT NO.
                      KIND
                      A1, 19950705,
PΙ
         R: CH, DE, FR, GB, IT, LICE CONTROL
     JP 07233131
                      A2 19950905
PRAI JP 1993-349879
                            19931229.
    MARPAT 123:340946
OS
GΙ
```

Amino acid esters of formula (XOA) nY [X = arom. or heterocyclic moiety; A AΒ = L-amino acid; n .gtoreq. 2; Y = moiety derived from compd. with 2 or more carbonyl or sulfonyl groups, bound to N of amino acid groups A] are claimed. The esters are hydrolyzed by leukocytes or elastase with high specificity, and are thus useful for detecting these in bodily fluids. For example, reaction of ClCO(CH2)4COCl (adipoyl dichloride) with excess L-alanine and pyridine gave 87.5% (CH2)4[CO-Ala-OH]2. This was treated with SOC12 to give its acid dichloride (100%), which reacted with 3-(dimethylamino)-5-piperidinophenol [prepn. given] to give 65.8% title compd. I. A test paper based on I and 1-diazo-2-naphthol-4-sulfonic acid gave color change (red-purple) for leukocyte liq. or elastase, but no change for .alpha.-chymotrypsin, trypsin, cathepsin, or butylcholinesterase, and only slight change for subtilisin. syntheses and a variety of diagnostic examples are provided.

ST amino acid ester detn leukocyte elastase

IT Diagnosis Leukocyte

> Urine (prepn. of amino acid esters as reagents for detecting leukocytes and elastase in bodily fluids)

IT Amino acids, preparation

RL: ARG (Analytical reagent use); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP हरेंद्र र एक क्राइट्टि (Preparation); USES (Uses)

(esters, prepn. of amino acid esters as reagents for detecting leukocytes and elastase in bodily fluids)

480-93-3P, 3-Hydroxyindole 13170-66-6P, Cyclohexane-1,4-dicarbonyl IT 136458-21-4P 141187-06-6P 170487-71-5P 40248-00-8P chloride 170487-74-8P 170487-75-9P 170487-73-7P 170487-72-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (intermediate; prepn. of amino acid esters as reagents for

detecting leukocytes and elastase in bodily fluids)

IT 9004-06-2, Elastase 🕒

· 24 - 1476年98 RL: ANT (Analyte); ANST (Analytical study)

(prepn. of amino acid esters as reagents for detecting leukocytes and elastase in bodily fluids)

170487-45-3P 170487-42-0P 170487-43-1P 170487-44-2P IT 170487-41-9P 170487-47-5P **170487-48-6P** 170487-49-7P 170487-46-4P

> Timothy Saunders EIC-LAW Lib. 308-413

·通知,对《美兴·魏 Action and property

主义 计图 化铁矿

```
170487-52-2P
                                                     170487-53-3P
                                                                     170487-54-4P
     170487-50-0P
                     170487-51-1P
                                     170487<u>-</u>57-7P
                                                     170487-58-8P
                                                                     170487-59-9P
     170487-55-5P
                     170487-56-6P
                                     170487-62-4P 170487-63-5P
     170487-60-2P
                     170487-61-3P
                                                                     170487-64-6P
                                     170487-67-9P
                     170487-66-8P
                                                     170487-68-0P
                                                                     170487-69-1P
     170487-65-7P
     170487-70-4P
     RL: ARG (Analytical reagent use); SPN (Synthetic preparation); THU
     (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (prepn. of amino acid esters as reagents for detecting
        leukocytes and elastase in bodily fluids)
                                                                        61 - 90 - 5,
     56-40-6, Glycine, reactions 56-41-7, L-Alanine, reactions
IT
     L-Leucine, reactions 72-18-4, L-Valine, reactions 83-72-7,
     2-Hydroxy-1,4-naphthoquinone 89-05-4, Benzene-1,2,4,5-tetracarboxylic
            89-32-7
                       99-14-9, Propane-1,2,3-tricarboxylic acid 100-20-9,
     1,4-Benzenedicarbonyl dichloride 108-73-6, 1,3,5-Trihydroxybenzene 110-89-4, Piperidine, reactions 111-50-2, Adipoyl chloride 119-80-2,
     2,2'-Dithiodibenzoic acid
                                  124-40-3, reactions
                                                         142-08-5,
                          533-75-5, Tropolone 585-47-7, Benzene-1,3-disulfonyl
     2-Hydroxypyridine
     chloride 608-08-2, Indoxyl acetate 636-78-2, 4-Sulfobenzoic acid
     1076-38-6, 4-Hydroxycoumarin 1076-97-7, 1,4-Cyclohexanedicarboxylic acid
     1141-38-4, Naphthalene-2,6-dicarboxylic acid 3119-64-0,
     Diphenylmethane-4,4'-disulfonyl chloride 3387-26-6, Furan-3,4-dicarboxylic acid 3739-94-4, Pyridine-2,6-dicarbonyl chloride
                 13827-62-8, Naphthalene-2,6-disulfonyl chloride dinicotinic acid 33177-29-6 67294-61-5,
                                                                       15658-35-2,
     4808-48-4
     6,6'-Dithiodinicotinic acid
     1,3,6-Naphthalenetrisulfonyl chloride 170487-76-0 170487-77-1
                                 170487-80-6
                    170487-79-3
     170487-78-2
     RL: RCT (Reactant)
        (starting material; prepn. of amino acid esters as reagents
        for detecting leukocytes and elastase; in bodily fluids)
IT
     170487-48-6P
     RL: ARG (Analytical reagent use); SPN (Synthetic preparation); THU
     (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (prepn. of amino acid esters as reagents for detecting
        leukocytes and elastase in bodily fluids)
RN
     170487-48-6 HCAPLUS
     L-Alanine, N,N'-[1,4-phenylenebis(oxycarbonyl)]bis-, bis[3-(dimethylamino)-
CN
     5-(1-piperidinyl)phenyl] ester (9CI) (CA INDEX NAME)
```

illist see of a fight

्यास्य स्वीत्रामा के स्वास्त्र स्वीत्रामा के

 $u^{*}I^{*}$ 

Absolute stereochemistry.

SEPREMITE

PAGE 1-B

L20 ANSWER 9 OF 12 HCAPLUS COPYRIGHT 2001 ACS

AN 1995:887807 HCAPLUS

DN 123:314522

TI Pharmaceutical composition for HIV protease inhibitor [ritonavir] with improved oral bioavailability

IN Al-razzak, Laman A.; Marsh, Kennan C.; Pyter, Richard A.

PA Abbott Laboratories, USA

SO PCT Int. Appl., 55 pp.

CODEN: PIXXD2

DT Patent

LA English

IC ICM A61K009-14

ICS A61K009-16; A61K009-48

CC 34-2 (Amino Acids, Peptides, and

Section cross-reference(s): 28, 63

11

FAN.CNT 2

PATENT NO. KIND DATE APPLICATION NO. DATE

e order or a second

```
PΙ
     WO 9509614
                       A1
                            19950413
                                               1994-US10096
         W: AU, CA, JP, KR
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                                            US 1994-297004
                            19960924
                                                              19940831
     US 5559158
                       Α
                       A1 # 199505019
                                           AU 1994-77229
                                                              19940909
     AU 9477229
                       B2
                            19980122
     AU 685509
                            19960717
                                            EP:1994-928043
                                                              19940909
                       A1 1
     EP 721330
                            20010328
     EP 721330
                       В1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
                                            JP 1994-510810
                                                              19940909
     JP 09503501
                       T2
                            19970408
                                            AT 1994-928043
                                                              19940909
     AT 200023
                       Ε
                            20010415
                            19931001
PRAI US 1993-130409
                       Α
     US 1994-267273
                            19940628
                       Α
                            19940831
     US 1994-297004
                       Α
                       W
                            19940909
     WO 1994-US10096
     MARPAT 123:314522
os
GΙ
```

11,

A solid pharmaceutical compn. is disclosed which comprises a AB pharmaceutically acceptable adsorbent or mixt. of adsorbents, to which is adsorbed a mixt. of: (1) a pharmaceutically acceptable org. solvent or mixt. of solvents; (2) an HIV protease-inhibiting compd.; and (3) one or more pharmaceutically acceptable acids. The solid compn. can optionally be encapsulated in a hard gelatin capsule. The compn. is particularly applicable to compd. I, and esp. its (2S,3S,5S,L)-isomer [ritonavir; II]. For example, oral administration of unformulated II to dogs gave < 2.0% mean bioavailability. In contrast, 89.6% mean bioavailability was obtained with the following capsule formulation: [II 21.84, propylene glycol 10.96, ethanol 22.99, Polysorbate 80 5.31, Cremophor EL 4.4, HCl 1.18, and Cab-o-sil 26.88% by wt. Also described are addnl. oral formulations (comparative and invention), and several syntheses of II. For example, N-(benzyloxycarbonyl) -L-phenylalaninol was converted in 5 steps to (2S,3S,5S)-PhCH2CH(NHZ)CH(OH)CH2CH(NHZ)CH2Ph [Z = benzyloxycarbonyl], which was deprotected and reacted with 5-thiazolylmethyl nitrophenyl carbonate to give intermediate III and its isomer from acylation of the other amino group. Coupling of III with N-[[N-methyl-N-[(2-isopropyl-4-thiazolyl)methyl]amino]carbonyl]-L-valine

```
[prepn. given] using the carbodiimide reagent EDC and
     1-hydroxybenzotriazole gave II.
     HIV protease inhibitor ritonavir pharmaceutical bioavailability; oral
ST
     bioavailability HIV protease inhibitor
     Drug bioavailability
ΙT
     Virucides and Virustats
         (pharmaceutical compn. of HIV protease inhibitor with improved oral
        bioavailability)
IT
     Castor oil
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (ethoxylated, Cremophor EL, adjuvant; pharmaceutical compn. of HIV
        protease inhibitor with improved oral bioavailability)
ΙT
     Pharmaceutical dosage forms
        (oral, pharmaceutical compn. of HIV protease inhibitor with improved
        oral bioavailability)
IT
     144114-21-6, Retropepsin
     RL: BSU (Biological study, unclassified); MSC (Miscellaneous); BIOL
     (Biological study)
         (HIV-1 and HIV-2; pharmaceutical compn. of HIV protease inhibitor with
        improved oral bioavailability)
     50-81-7, Ascorbic acid, biological studies 57-55-6, Propylene glycol,
IT
     biological studies
                           64-17-5, Ethanol, biological studies
     Citric acid, biological studies 104-15-4, p-Toluenesulfonic acid,
     biological studies 7631-86-9, Silica, biological studies 7647-01-0, Hydrochloric acid, biological studies 9002-96-4 9004-34-6, Cellulose,
                            9005-25-8, Corn starch, biological studies
     biological studies
                                   9050-36-6, Maltodextrin 14807-96-6, Talc,
     9005-65-6, Polysorbate 80
                            41080-67-5
                                             1.5
     biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
     (adjuvant; pharmaceutical compn. of HIV protease inhibitor with improved oral bioavailability) (115-08-2P, Thioformamide 13515-65-6P, 2-Methylpropanethioamide
IT
     32955-21-8P, 2-Amino-5-(ethoxycarbonyl)thiazole 32955-22-9P, Ethyl
     thiazole-5-carboxylate 33142-21-1P, Ethyl 2-chloro-2-formylacetate
     38585-74-9P, 5-(Hydroxymethyl)thiazole 59830-60-3P, N-[[(Benzyl)oxy]carbonyl]-L-phenylalaninal 65386-28-9P,
     4-(Chloromethyl)-2-isopropylthiazole hydrochloride
                                                             137649-69-5P
                   144141<del>-</del>68-4P
     143838-10-2P
                                     154212-60-9P,
                                                      154212-59-6P
     144163-97-3P
                     144164-10-3P
                                     144164-11-4P
     2-Isopropyl-4-[(N-methylamino)methyl]thiazole
                                                        154212-61-0P
     154248-99-4P 162537-10-2P
                                   162849-92-5P
                                                   162849-93-6P
                                    162849-96-9P 162990-03-6P
                     162849-95-8P
     162849-94-7P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (intermediate; pharmaceutical compni of HIV protease inhibitor with
        improved oral bioavailability)
                     162990-01-4P
IT
     155213-67-5P
     RL: BPR (Biological process); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC
     (Process); USES (Uses)
         (pharmaceutical compn. of HIV protease inhibitor with improved oral
                                     ONE OF BUILDING
        bioavailability)
     62-56-6, Thiourea, reactions 74-89-5, Methylamine, reactions
IT
     Formamide, reactions 98-80-6, Phenylboronic acid 105-39-5, Ethyl chloroacetate 109-94-4, Ethyl formate 3534-07-6, 1,3-Dichloroacetone
     563-83-7, Isobutyramide 6306-52-1, L-Valine methyl ester hydrochloride 6372-14-1, N-[[(Benzyl)oxy]carbonyl] L-phenylalaninol 7693-46-1,
     4-Nitrophenyl chloroformate 24424-99-5, Di-tert-butyl dicarbonate
     40635-67-4, .alpha.-Acetoxyisobutyryl bromide 126147-70-4,
     N-(Phenoxycarbonyl)-L-valine 156732-13-7 156732-15-9
                                                                     169597-13-1
                                     RL: RCT (Reactant)
                                     High the party
```

(starting material; pharmaceutical compn. of HIV protease inhibitor with improved oral bioavailability)

IT 162537-10-2P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (intermediate; pharmaceutical compn. of HIV protease inhibitor with improved oral bioavailability)

162537-10-2 HCAPLUS RN

L-Valine, N-[(4-nitrophenoxy)carbonyl] + methyl ester (9CI) CN (CA INDEX

Absolute stereochemistry.

126147-70-4, N-(Phenoxycarbonyl)-L-valine RI: RCT (Reactant) IT

(starting material; pharmaceutical compn. of HIV protease inhibitor with improved oral bioavailability)

126147-70-4 HCAPLUS RN

1993年第4月 L-Valine, N-(phenoxycarbonyl) - (9GI) (CA INDEX NAME) CN

Absolute stereochemistry.

L20 ANSWER 10 OF 12 HCAPLUS COPYRIGHT 2001 ACS

AN 1990:179761 HCAPLUS

DN 112:179761

Optical resolution of amino acid derivatives by high-performance liquid TI chromatography on tris(phenylcarbamate)spof cellulose and amylose

ΑU Okamoto, Yoshio; Kaida, Yuriko; Aburatani, Ryo; Hatada, Koichi

CS

1 1

Fac. Eng. Sci., Osaka Univ., Toyonaka, 560, Japan J. Chromatogr. (1989), 477(2), 367-76 CODEN: JOCRAM; ISSN: 0021-9673

DTJournal

SO

English LΑ

34-2 (Amino Acids, Peptides, and Ario CC 事"新兴" Proteins)

AB The optical resoln. of 10 N-protected alanine esters was examd. by HPLC using 6 cellulose and 5 amylose tris(phenylcarbamate) derivs. as chiral stationary phases. Tris(3,5-dimethylphenylcarbamate)s of both cellulose and amylose showed high resolving power for these racemates. The resoln. of 23 N-benzyloxycarbonyl .alpha.-amino acid ester was also tested on tris(3,5-dimethylphenylcarbamate)s of cellulose and amylose. All but 2 amino acid derivs. were completely resolved at least by one of the columns. On cellulose tris(3,5-dimethylphenylcarbamate), all L-amino

```
acids (except threonine) eluted first.
     resoln amino ester HPLC cellulose trisphenylcarbamate; amylose
ST
     trisphenylcarbamate resoln amino ester
     Resolution
IT
         (of protected amino acid esters by HPLC on cellulose and
        amylose tris(phenylcabamate) stationary phases)
     Chromatography, column and liquid
IT
         (high-performance, resoln. by, of protected amino acid esters with
        cellulose and amylose tris(phenylcarbmate) stationary phases)
IT
     Amino acids, esters
     RL: PROC (Process)
         (N-protected, esters, resoln. of, by HPLC with cellulose and
        amylose tris(phenylcarbamate): stationary, phases)
                               5446-46-8, N-Benzoyl-DL-alanine ethyl ester
IT
                   5143-72-6
     5513-39-3, N-Benzyloxycarbonyl-DL-alanine benzyl ester
                                                                    5557-84-6
     23161-27-5, N-Benzyloxycarbonyl-DL-serine ethyl ester
     N-Benzyloxycarbonyl-DL-threonine ethyl ester
                                                         25282-53-5,
     DL-Phenylalanine benzyl ester 39978-35-3, N-Benzyloxycarbonyl-DL-leucine
                     40489-45-0, N-Benzyloxycarbonyl-DL-serine benzyl ester
     ethyl ester
     42998-42-5, N-Benzyloxycarbonyl-DL-valine ethyl ester
                                                                    42998-44-7,
     N-Benzyloxycarbonyl-DL-phenylalanine ethyl ester
                                                             42998-45-8
     46229-47-4, DL-Alanine benzyl ester 72604-32-1, N-tert-Butoxycarbonyl-DL-
     alanine ethyl ester # 72604-33-2; N-Benzyloxycarbonyl-DL-alanine ethyl
              85369-24-0 86827-19-2 88406-41-1
                                                      103063-37-2,
     ester
     N-Benzyloxycarbonyl-DL-phenylalanine benzyl ester 11
N-Acetyl-DL-alanine benzyl ester 126400-85-9 126400
126400-88-2 126400-89-3 126400-90-6 126400-91-7
                                                               114285-13-1,
                                                            126400-86-0
                                                                            126400-87-1
                                                                    126400-92-8
                     126400-94-0, N-Benzoyl-DL-alanine benzyl ester
     126400-93-9
                    126400-97-3, N-Benzyloxycarbonyl-DL-valine benzyl
     126400-96-2
              126400-98-4, N-Benzyloxycarbonyl-DL-norvaline ethyl ester
     126400-99-5, N-Benzyloxycarbonyl-DL-norvaline benzyl ester
                                                                        126401-00-1,
     N-Benzyloxycarbonyl-DL-leucine benzyl ester
                                                        126401-01-2,
                                                           126401-02-3,
     N-Benzyloxycarbonyl-DL-norleucine ethyl ester
     N-Benzyloxycarbonyl-DL-norleucine benzyl ester 126401-03-4,
     N-Benzyloxycarbonyl-DL-isoleucine benzyl ester
                                                            126401-04-5
                                                                            126401-05-6
     126401-06-7, N-Benzyloxycarbonyl-DL-threonine benzyl ester
                                                                         126401-07-8
                                    126401-10-3 126401-11-4
                                                                    126401-12-5
     126401-08-9
                     126401-09-0
                                    126401-15-8 126401-16-9
126401-20-5 126401-21-6,
                     126401-14-7
                                                                    126401-17-0
     126401-13-6
     126401-18-1
                    126401-19-2
     N-Benzyloxycarbonyl-DL-methionine benzyliester
                                                            126401-22-7
                                                                            126401-23-8
     126433-60-1, N-Benzyloxycarbonyl-DL-methionine ethyl ester
                    126456-13-1, N-Benzyloxycarbonyl-DL-proline ethyl ester
     126456-12-0
     126456-14-2, N-Benzyloxycarbonyl-DL-proline benzyl ester
     RL: PROC (Process)
         (resoln. of, by HPLC with cellulose and amylose
         tris(phenylcarbamate) stationary phases)
     9047-05-6, Amylose tris(phenylcarbamate) 4 9047-07-8, Cellulose
IT
     tris (phenylcarbamate) 103938-40-5, Cellulose tris (4-bromophenylcarbamate) 103938-45-0, Cellulose tris (4-ethylphenylcarbamate) 103938-46-1, Cellulose tris (4-fluorophenylcarbamate) 103938-49-4 107028-63-7, Amylose tris (3,5-dichlorophenylcarbamate) 115901-94-5, Amylose
     tris(4-chlorophenylcarbamate) 115901-96-7, Amylose tris(4-
                                        and the first of the second
     methylphenylcarbamate)
                                      好的 维西拉 化性
     RL: RCT (Reactant)
         (stationary phase, for HPLC resoln, of protected alanine
                                      主义引发控制 清净和公司
     103938-44-9, Cellulose tris(3,5-dimethylphenylcarbamate)
                                                                       112049-40-8
IT
     RL: RCT (Reactant)
         (stationary phase, for HPLC resoln, of protected amino acid
```

- 1 to the second secon

```
esters)
ΙT
    88406-41-1 126400-96-2
    RL: PROC (Process)
        (resoln. of, by HPLC with cellulose and amylose
       tris(phenylcarbamate) stationary phases)
    88406-41-1 HCAPLUS
Alanine, N-(phenoxycarbonyl)-, ethyl ester (9CI) (CA INDEX NAME)
RN
CN
    0
Pho-c-NH
   Me-CH-C-OEt
    126400-96-2 HCAPLUS
RN
                                  phenylmethyl ester (9CI) (CA INDEX NAME)
CN
    Alanine, N-(phenoxycarbonyl)-,
            NH-
Ph-CH2-O-C-CH-Me
L20
    ANSWER 11 OF 12
                     HCAPLUS
                              COPYRIGHT
    1988:611486 HCAPLUS
AN
DN
     109:211486
    Preparation of heterocyclylpeptides as renin inhibitors
ΤI
    Ryono, Denis Evan; Weller, Harold Norris, III
IN
PA
SO
    CODEN: EPXXDW
DT
    Patent
    English
LΑ
    ICM C07K005-06
IC
     ICS C07K005-08; C07D233-64; C07D417-12; C07D401-12; C07D403-12;
         A61K037-02; A61K037-64; A61K031-415; A61K031-425; A61K031-44
    C07D417-12, C07D277-00, C07D233-00; C07D401-12, C07D213-00, C07D233-00;
     C07D417-12, C07D235-00, C07D233-00
CC
     34-3 (Amino Acids, Peptides, and
    Proteins)
     Section cross-reference(s): 1
FAN.CNT 1
                                          APPLICATION NO.
     PATENT NO.
                     KIND
                           DATE
                           19870812 EP 1987-101373
                      A2
                                                           19870202
ΡI
    EP 231919
                           19900718
    EP 231919
                      A3
    EP 231919
                      B1
                           19930120
        R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE
                           19870930 ZA 1987-563
                                                          19870126
     ZA 8700563
                           19870806
                      A1:
                                         AU 1987-68153
                                                          19870130
    AU 8768153
    AU 595578
                      B2
                           19900405
                                         DK 1987-523
                                                           19870202
    DK 8700523
                      A ·
                           19870804
                           19880328
                                          HU 1987-352
                                                           19870202
    HU 44576
                      A2
    HU 203368
                      В
                           19910729
```

州州中北市情

AT: 1987-101373

19870202

E.

1. . [1]

19930215

AT 84793

```
Т3
                                                                       ES 1987-101373
                                                19940101
                                                                                                        19870202
        ES 2043611
                                       A2 19871110 JP 1987-23434
A1 19921124 CA 1987-528852
A 19891205 US 1989-373633
19860203
                                                                                                        19870203
        JP 62258365
        CA 1310792
                                                                                                        19870203
                                                                                                        19890629
        US 4885292
PRAI US 1986-825724
        US 1987-3446
                                                19870115
        EP 1987-101373
                                                19870202
        x-(NHCHR5CO)p-NHCHR4CONHCHR3CH(OH)R1 [I; x=R6(CH2)n, R6(CH2)nO2C,
AB
        R60(CH2)nCO, R6(CH2)nSO2, etc.; R1 = (substituted) N-contg. heterocyclyl;
        R3, R4, R5 = (halo)alkyl, arylalkyl, hydroxyalkyl, carboxyalkyl,
        guanidinoalkyl, imidazolylalkyl, etc.; R6 = alkyl, cycloalkyl, aryl,
        heterocyclyl; n = 0-5; p = 0, 1] useful as renin inhibitors, (no data)
        were prepd. N-[(1,1-Dimethylethoxy)carbonyl]-L-leucinal (prepn. from
        leucine given) in THF was added to a mixt. of BuLi and
        1-[(phenylmethoxy)methyl]-1H-imidazole in THF at -70.degree.. The mixt.
        was kept at -70.degree. for 1 h, and at 0.degree. for 15 min followed by
        quenching with aq. NH4Cl and hydrolysis with HCl/EtOAc to give
        .alpha.-[(S)-1-amino-3-methylbutyl]-1-[(phenylmethoxy)methyl]-1H-imidazole-
        2-methanol-2HCl. The latter was coupled with N-[N-[(1,1-
        dimethylethoxy)carbonyl]-L-phenylalanyl]-l'-[phenylmethoxy)methyl]-L-
        histidine (prepn. given) using 1-hydroxybenzotriazole and DCC in DMF at
        ice temps. The coupling product was hydrogenolyzed in aq. MeOH contg. HCl
        over Pd/C to give N2-[N-[(1,1-dimethylethoxy)carbonyl]-L-phenylalanyl]-N-
        [(S)-1-[hydroxy(1H-imidazol-2-yl)methyl]-3-methylbutyl]-L-histidinamide-
        imidazolylpeptide prepn antihypertensive; Renin inhibitor peptide
ST
        heterocyclyl; antihypertensive heterocyclylpeptide
IT
        Antihypertensives
              (heterocyclyl-contg. peptide alcs.)
        Peptides, preparation for the first the periods
IT
        RL: SPN (Synthetic preparation); PREP (Preparation)
              (heterocyclyl-contg., prepn. of, as renin inhibitors)
IT
        7693-46-1, 4-Nitrophenyl chloroformate
        RL: RCT (Reactant)
                                                          4 50 新 1 4
              (acylation by, of phenylalanine Me ester, in prepn. of
                                                            antihypertensive) 🕌
        123-75-1, Pyrrolidine, reactions in the Prince
IT
        RL: RCT (Reactant)
                                                           · 建氯基苯基基苯基基基
              (acylation of, by phosgene, in prepn. of antihypertensive)
                                                           THE COURT OF THE PARTY OF THE P
        100-39-0, Benzyl bromide
ΙT
        RL: RCT (Reactant)
              (alkylation by, of di-Et benzylmalonate, in prepn. of antihypertensive)
        607-81-8, Diethyl benzylmalonate
ΙT
        RL: RCT (Reactant)
                                          . -1
              (alkylation of, by benzyl bromide)
        105-53-3, Diethyl malonate
ΙT
        RL: RCT (Reactant)
              (alkylation of, by chloromethylnaphthalene, in prepn. of
              antihypertensive)
ΙT
        6638-79-5, O,N-Dimethylhydroxylamine hydrochloride
        RL: RCT (Reactant)
              (amidation by, of (dimethylethoxycarbonylamino) cyclohexanepropanoic
              acid, in prepn. of antihypertensive)
        7524-50-7, L-Phenylalanine methyl ester hydrochloride
IT
                                                           11 1 1 1 1 1
        RL: RCT (Reactant)
              (amidation by, of pyrrolidinecarbonyl chloride, in prepn. of
              antihypertensive)
IT
        13139-15-6
        RL: RCT (Reactant)
              (amidation of, by dimethylhydroxylamine; in prepn. of antihypertensive)
```

Tary Milaine

14. 为为 ...

```
86-52-2, 1-Chloromethylnaphthalene
IT
     RL: RCT (Reactant)
        (condensation of, with di-Et malonate, in prepn. of antihypertensive)
     61-90-5, L-Leucine, reactions
IT
     RL: RCT (Reactant)
        (conversion of, to Et ester)
     71-00-1, L-Histidine, reactions
IT
     RL: PROC (Process)
        (conversion of, to Me ester, in prepn. of antihypertensive)
                                         42 pt 1942 1
IT
     115888-02-3P
     RL: SPN (Synthetic preparation); FORM (Formation, nonpreparative); PREP
        (formation of, in prepn. of antihypertensive)
                          49822-58-4
     288-47-1, Thiazole
IT
     RL: RCT (Reactant)
        (lithiation and condensation of, with leucinal deriv., in prepn. of
        antihypertensive)
     623-33-6, Glycine ethyl ester monohydrochloride 2666-93-5, L-Leucine
IT
                                 25024-53-7 25616-33-5
                    13734-34-4
    methyl ester
                  96867-10-6
     83468-83-1
     RL: RCT (Reactant)
     (peptide coupling of, in prepn. of antihypertensive) 7389-87-9P, L-Histidine methyl ester dihydrochloride
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and N-protection of, in prepn. of antihypertensive)
     111629-38-0P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and coupling of, with phenylalanine deriv., in prepn. of
        antihypertensive)
                                  一人, 他严惧的
                 115766-40-0P
IT
     4372-32-1P
                                              . .
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and decarboxylation of in prepn. of antihypertensive)
                    115766-27-3P
ΙT
     115766-26-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and deprotection of in prepn. of antihypertensive)
IT
     82010-31-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and oxidn. of, in prepn. of antihypertensive)
                    115766-29-5P 115766-31-9P 115766-34-2P 115766-35-3P 115766-38-6P
IT
     115766-28-4P
     115766-37-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and peptide coupling of, in prepn. of antihypertensive) 3-40-6P
IT
     7533-40-6P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and protection of, in prepn. of antihypertensive)
                  111629-37-9P 115766-33-1P 115766-41-1P.
IT
     20898-43-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and reaction of, in prepn. of antihypertensive) 3-40-0P
IT
     2743-40-0P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and redn. of)
     51987-73-6P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and redn. of, in prepn. of antihypertensive)
629-39-1P
                                        学规体(1987)
     111629-39-1P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and sapon. of, in prepn. of antihypertensive)
                                   115765-67-8P 115765-68-9P 115765-69-0P
                    115765-66-7P
IT
     115765-65-6P
                    115765-71-4P 115765-72-5P 115765-73-6P 115765-74-7P
     115765-70-3P
                                   115765-78-1P 115765-79-2P
                                                                  115765-80-5P
     115765-76-9P
                   115765-77-0P
```

(有) 有针 等特 (17.5 ) (1.5 ) (17.5 ) (17.5 ) (17.5 )

```
115765-81-6P
                    115765-82-7P
                                   115765-83-8P
                                                  115765-84-9P
                                                                  115765-86-1P
                                                115802-91-0P
    115765-88-3P
                    115766-42-2P
                                   115766-43-3P
                                               o d y
                                                                  115887-98-4P
                    115888-04-5P
                                   115888-05-6P
                                                  115888-06-7P:
                                                                 `115888-07-8P
    115888-03-4P
                                   115888-86-3P
                    115888-09-0P
                                                  115935-94-9P
                                                                 115935-95-0P
     115888-08-9P
     115935-96-1P
    RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic
    preparation); BIOL (Biological study); PREP (Preparation)
        (prepn. of, as antihypertensive) 30189-51-6P 37736-82-6P
IT
     597-55-7P
                                              66605-57-0P
                   58521-45-2P
                                 64152-76-7P
     54601-21-7P
                                                             102639-05-4P
     72155-45-4P
                   87694-50-6P
                                 98105-42-1P
                                               102639-04-3P
                                   110695-91-5P
                    104539-18-6P
                                                  114457-62-4P
                                                                  114473-20-0P
     103322-56-1P
                                   115765-91-8P 115765-92-9P
                                                                  115765-93-0P
                    115765-90-7P
     115765-89-4P
                                  115765-96-3P
                    115765-95-2P
                                                  115765-97-4P
                                                                  115765-98-5P
     115765-94-1P
                                                  115766-02-4P
                                                                  115766-03-5P
                    115766-00-2P
                                   115766-01-3P
     115765-99-6P
                                   115766-06-8P
                                                  115766-07-9P
                                                                  115766-08-0P
                    115766-05-7P
     115766-04-6P
                    115766-10-4P
                                   115766-11-5P
                                                  115766-12-6P
                                                                  115766-13-7P
     115766-09-1P
     115766-14-8P
                    115766-15-9P
                                   115766-16-0P
                                                  115766-17-1P
                                                                  115766-18-2P
                                                  115766-23-9P
                                                                  115766-24-0P
     115766-19-3P
                    115766-20-6P
                                   115766-22-8P
                                                  115766-39-7P
                                                                  115802-92-1P
                    115766-32-0P
                                   115766-36-4P
     115766-25-1P
                    115802-94-3P
                                   115802-95-4P . 115887-99-5P
                                                                  115888-00-1P
     115802-93-2P
                                    長記したのよう性 不良
     115888-01-2P
    RL: SPN (Synthetic preparation) PREP (Preparation)
        (prepn. of, as intermediate for antihypertensive) 766-30-8P
IT
     115766-30-8P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of, as intermediates for antihypertensive)
     1070-83-3, 3,3-Dimethylbutanoic acid
IT
     RL: RCT (Reactant)
                                  (reaction of, in prepn. of antihypertensive peptide deriv.)
     109-01-3, 1-Methylpiperazine *110-91-8, Morpholine, reactions
IT
                                  国籍对"国"的原则,国
     RL: RCT (Reactant)
        (reaction of, with (nitrophenoxycarbonyl)phenylalanine)
     70-34-8, 2,4-Dinitrofluorobenzene
IT
     RL: RCT (Reactant)
        (reaction of, with histidine contg. dipeptide, in prepn. of
                                  g Marie Con
        antihypertensive)
     68-11-1, Mercaptoacetic acid, uses and miscellaneous
ΙT
                                  147,20万亩114.7
     RL: USES (Uses) 🖟 📑
        (reagent, for deprotection of dinitrophenylhistidine contg.
                                      10 · 16
        peptide)
ΙT
     3674-06-4
     RL: RCT (Reactant)
        (redn. of, in preph. of antihypertensive)
                                       医原种维斯 中一
IT
     54601-21-7P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of, as intermediate for antihypertensive) 01-21-7 HCAPLUS
     54601-21-7 HCAPLUS
RN
CN
     L-Phenylalanine, N-['(4-nitrophenoxy)carbonyl];, methyl ester (9CI)
                                    (tajara)
     INDEX NAME)
```

Absolute stereochemistry.

11:

```
\begin{array}{c|c} O & H & O \\ \hline O & N & S \\ \hline O & O \\ \hline O & Ph \\ \end{array}
```

```
COPYRIGHT 2001
     ANSWER 12 OF 12
                      HCAPLUS
     1974:437818 HCAPLUS
AN
     81:37818
DN
ΤI
     Histidine derivatives
     Tanabe Seiyaku Co., Ltd.
PA
SO
     Brit., 9 pp.
     CODEN: BRXXAA
DT
     Patent
LΑ
     English
IC
     C07D; A61K
CC
     34-3 (Synthesis of Amino Acids, Peptides
     and Proteins)
FAN.CNT 1
                      KIND
     PATENT NO.
                             19740508 GB 1971-12019
     GB 1352414
                       Α
                                                              19710428
PΙ
     For diagram(s), see printed CA Issue.
GΙ
     L-Pyroglutamyl-L-proline amide (I), a thyrotropine-releasing or thyroid
AB
     gland-stimulating factor, was prepd. from L-histidine Me ester (II).
     III-IV were prepd. from II by successive treatment with the appropriate
     chloride or azidoformate, hydrolysis, and tosylation. Condensation of these with L-proline amide at -10 to 10 degree. in the presence of a
     carbodiimide reagent, followed by treatment with
     N-benzyloxycarbonyl-L-pyroglutamickacid, and removal of the protecting
     groups, gave I.
                                 · 持續對理性表現了 1
ST
     histidine deriv
IT
     Protective groups
        (alkoxycarbonyl and sulfenyl groups, for amino group of histidine
        methyl ester, in peptide prepn.)
IT
     Peptides, preparation
     RL: PREP (Preparation)
        (o-Nitrophenyl) sulfenyl group
IT
     (tert-Pentyloxy) carbonyl group
     [(p-Methoxybenzyl)oxy]carbonyl group
                                              1.12
     tert-Butoxycarbonyl group
        (protective group, for amino group in peptide prepn.)
IT
                                  Amino group
        (protective groups for, alkoxycarbonyl and sulfenyl groups as, in
                                    四种 计
        peptide prepn.)
                                    Mr Leiter W
IT
     Imido group
        (protective groups for, benzyl and tert-akyloxycarbonyl groups as, in peptide prepn.)
nzyloxy) carbonyl group
IT
     (Benzyloxy) carbonyl group
        (protective groups, for imido group in peptide prepn.)
IT
     7531-52-4
     RL: RCT (Reactant)
        (condensation of, with histidines)
                                                               35899-43-5P
                                  23241-51-2P 24305-27-9P
                   23241-50-1P
IT
     15354-08-2P
                                                35899-47-9P
                                                               35899-49-1P
                                  35899-46-8P
     35899-44-6P
                   35899-45-7P
```

1 11 1 11

1 1.1

ii.

```
53090-21-4P
     53090-17-8P 53090-19-0P
                                             53090-24
     RL: SPN (Synthetic preparation) PREP (Preparation)
        (prepn. of)
IT
     98-79-3
                                  ,在教育为2011集合
     RL: RCT (Reactant)
        (protective group for imido group of, in peptide prepn.)
IT
     7389-87-9
     RL: RCT (Reactant)
        (protective groups for .alpha -amino group of, in peptide prepn.)
                             24608-52-4 40438-33-3
                 7669-54-7
IT
     5591-72-0
     RL: RCT (Reactant)
                                             11:1
        (reaction of, with histidine methyl ester)
                  53090-23-6
IT
     32159-21-0
     RL: RCT (Reactant)
        (reaction of, with histidyl prolinamide)
ΙT
     53090-17-8P 53090-19-0P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
     53090-17-8 HCAPLUS
RN
     L-Histidine, N-[(4-methoxyphenoxy)carbonyl]-, methyl ester (9CI) (CA
CN
     INDEX NAME)
```

Absolute stereochemistry

RN 53090-19-0 HCAPLUS

CN L-Histidine, N-[(4-methoxyphenoxy)carbonyl]-1-[(4-methylphenyl)sulfonyl]-,
compd. with N-cyclohexylcyclohexanamine (1:1) (9CI) (CA INDEX NAME)

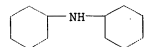
CM 1

CRN 53090-18-9 CMF C21 H21 N3 O7 S CDES 5:L

Absolute stereochemistry.

CM 2

CRN 101-83-7 CMF C12 H23 N



L1 STR

NODE ATTRIBUTES:

HCOUNT	IS	M1	AΤ	10	
NSPEC	IS	R	ΑT	1	
NSPEC	IS	R	AΤ	2	
NSPEC	IS	R	ΑT	3	
NSPEC	IS	R	AΤ	4	
NSPEC	IS	R	AT	5	
NSPEC	IS	R	AΤ	6	i
NSPEC	IS	С	ΑT	7	
NSPEC	IS	С	ΑT	8	
NSPEC	IS	С	ΑT	9	
NSPEC	IS	С	ΑT	10	ŀ
NSPEC	IS	С	AT	11	
NSPEC	IS	C	AΤ	12	
NSPEC	IS	С	AT	13	
NSPEC	IS	С	AT	14	
DEFAULT	MLI	EVEL IS	ATO	M	
	T (4	GT 3 G G	200	7	

MLEVEL IS CLASS AT 7 8 9 10 11 12 13 1

DEFAULT ECLEVEL IS LIMITED

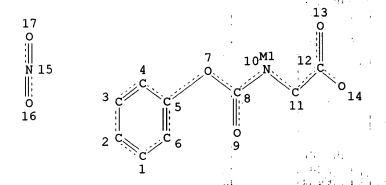
# GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

SILL	TO WILKIDOL	25. HOME						
L3	501	SEA FILE=REGISTRY SSS FUL L1						
L4	213	SEA FILE=HCAPLUS ABB=ON PLU=ON L3						
L6		SEA FILE=HCAPLUS ABB=ON PLU=ON 2000:755247/AN						
L14	75483	SEA FILE=HCAPLUS ABB=ON PLU=ON (AMINO ACIDS OR PEPTIDES OR						
		PROTEINS)/CC						
L15	83	SEA FILE=HCAPLUS ABB=ON PLU=ON L4 AND L14						
L16	82	SEA FILE=HCAPLUS ABB=ON PLU=ON L15 NOT L6						
L18	4	SEA FILE=HCAPLUS ABB=ON PLU=ON (HPLC OR CHROMATOGARPH?) AND						
		L16						
L19	9	SEA FILE=HCAPLUS ABB=ON PLU=ON, (ENANTIO? OR SEPARAT? OR						
CHROMOPHO? OR REAGENT) AND L16								
L20	12	SEA FILE=HCAPLUS ABB=ON PLU=ON L18 OR L19						
L21		STR						



NODE ATT	rri	BUTES:			
HCOUNT	IS	M1	AΤ	10	
NSPEC	IS	R	AΤ	1	
NSPEC	IS	R	AΤ	2	į:
NSPEC	IS	R	AΤ	3	1,
NSPEC	IS	R	AΤ	4	
NSPEC	IS	R	AΤ	5	
NSPEC	IS	R	AΤ	6	
NSPEC	IS	C	AΤ	7	
NSPEC	IS	C	AΤ	8	
NSPEC	IS	С	AΤ	9	
NSPEC	IS	С	AΤ	10	
NSPEC	IS	С	AΤ	11	
NSPEC	IS	С	AΤ	12	
NSPEC	IS	С	AT	13	•
NSPEC	IS	С	AT	14	
NSPEC	IS	С	AΤ	15	
NSPEC	IS	С	AΤ	16	
NSPEC	IS	С	AΤ	17	
DEFAULT	ML	EVEL IS	ATO	M	
				_	10.3

MLEVEL IS CLASS AT 7 8 9 10 11 12 13 14 15 16 17
DEFAULT ECLEVEL IS LIMITED

# GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 17

## STEREO ATTRIBUTES: NONE

L23	87	SEA	FILE=REGISTRY	Y SUB=L3	SSS FUL	L21		
L24	56	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	L23		
L25	50	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	L24	TON	L20
L26	49	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	L25	NOT	L6
L27	24	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	L14	AND	L26

```
ANSWER 1 OF 24 HCAPLUS
                                  COPYRIGHT 2001 ACS
L27
     2001:416903 HCAPLUS
AN
DN
     135:33643
     Preparation of 3-(2-aminoethylthio)methyl-4-oxo-4-(3-pyridyl)butanoic acid
ΤI
     derivatives as neuroprotective agents
     Bhagwat, Shripad; Palanki, Moorthy; Erdman, Paul; Doubleday, Mary; Sato,
IN
     Nippon Kayaku Co., Ltd., Japan
PA
so
     PCT Int. Appl., 119 pp.
     CODEN: PIXXD2
DT
     Patent
LΑ
     English
IC
     ICM C07D213-55
     ICS C07D213-56; C07D417-12; A61K031-44; A61P025-00
CC
      34-2 (Amino Acids, Peptides, and
      Section cross-reference(s): 1, 27
FAN.CNT 1
     PATENT NO.
                         KIND
                                DATE
                                                 APPLICATION NO.
PΙ
     WO 2001040187
                          A2
                                20010607
                                                WO 2000-JP8090
              AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
               CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
              HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU,
               LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD,
               SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA,
               ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, EI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRAI US 1999-450245
                                19991129
GΙ
```

 $R^{1}$ 

The title compds. [I; R1 = (un) substituted alkyl, aryl, arylakyl, etc.; R2 = OR2a, NR2bR2c; R3 = H, :O, CO2R3a, etc.; R4 = H, (un) substituted alkyl, aryl, etc.; R3 and R4 taken together = (un) substituted heterocyclyl; R5 = H, (un) substituted alkyl; R2a = H, (un) substituted alkyl, aryl, etc.; R2b, R2c = H, (un) substituted alkyl, aryl, etc.; NR2bR2c = (un) substituted heterocyclyl; R3a = H, (un) substituted alkyl, aryl, etc.] which have

till.

ind) The

```
utility in the treatment of conditions which benefit from administration
of neuroprotective agents generally, including treatment of central and
peripheral nervous condition as well as for promoting nerve cell
differentiation, were prepd. Thus, reacting 4-oxo-3-(piperidylmethyl)-4-
(3-pyridyl)butanoic acid with Me N-acetyl-L-cysteine ester in EtOH
afforded 85% (R)-II. Biol. data for compds. I were given.
aminoethylthiomethyloxopyridylbutanoic acid prepn neuroprotectant;
pyridylbutanoic acid aminoethylthiomethyloxo prepn neuroprotectant
                               The first of
Cytoprotective agents
   (neuroprotectants; prepn. of 3-(2-aminoethylthio)methyl-4-oxo-4-(3-
   pyridyl)butanoic acid derivs. as neuroprotective agents)
343574-29-8P
               343574-71-0P
                               343575-87-1P
                                              343575-92-8P
                                                              343577-16-2P
               343577-23-1P
                              343577-91-3P
343577-17-3P
RL: BAC (Biological activity or effector, except adverse); RCT (Reactant);
SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
study); PREP (Preparation); USES (Uses)
   (prepn. of 3-(2-aminoethylthio)methyl-4-oxo-4-(3-pyridyl)butanoic acid
   derivs. as neuroprotective agents)
220301-04-2P
               343573-75-1P
                               343573-76-2P
                                              343573-77-3P
                                                              343573-78-4P
343573-79-5P
               343573-80-8P
                              343573-81-9P
                                              343573-82-0P
                                                              343573-83-1P
                               343573-86-4P
343573-84-2P
               343573-85-3P
                                              343573-87-5P
                                                              343573-88-6P
                              343573#91-1P 343573-92-2P
343573-89-7P
               343573-90-0P
                                                             :343573-93-3P
                              1343573-96-6P 🐬 343573-97-7P
343573-94-4P
               343573-95-5P
                                                              343573-98-8P
343573-99-9P
               343574-00-5P
                              +343574-01-6P
                                              343574-02-7P:
                                                              343574-03-8P
                               343574-06-1P 343574-07-2P
343574-04-9P
               343574-05-0P
                                                              343574-08-3P
                              343574-11-8P
              343574-10-7P
                                                              343574-13-0P
343574-09-4P
                                              343574-12-9P
343574-14-1P
              343574-15-2P
                              343574-16-3P 343574-17-4P
                                                              343574-18-5P
                               343574-21-0P
343574-19-6P
               343574-20-9P
                                              343574-22-1P
                                                              343574-23-2P
343574-24-3P
               343574-25-4P
                              343574-26-5P
                                              343574-27-6P
                                                              343574-28-7P
                               343574-32-3P
                                              343574-33-4P
343574-30-1P
               343574÷31-2P
                                                              343574-34-5P
343574-35-6P
               343574-36-7P
                               343574-37-8P
                                              343574-38-9P
                                                              343574-39-0P
343574-40-3P
               343574-41-4P
                               343574-42-5P
                                              343574-43-6P
                                                              343574-44-7P
343574-45-8P
               343574-46-9P
                               343574-47-0P
                                              343574-48-1P
                                                              343574-49-2P
               343574-51-6P
                               343574-52-7P
                                              343574-53-8P
                                                              343574-54-9P
343574-50-5P
                              343574-57-2P
343574-55-0P
               343574-56-1P
                                              343574-58-3P
                                                              343574-59-4P
343574-60-7P
               343574-61-8P
                               343574-62-9P
                                              343574-63-0P
                                                              343574-64-1P
343574-65-2P
               343574-66-3P
                              343574-67-4P
                                              343574-68-5P
                                                              343574-69-6P
343574-70-9P
               343574-72-1P
                              343574-73-2P
                                              343574-74-3P
                                                              343574-75-4P
343574-76-5P
               343574-77-6P
                              343574-78-7P
                                              343574-79-8P
                                                              343574-80-1P
               343574-82-3P
                              343574-83-4P
                                              343574-84-5P
343574-81-2P
                                                              343574-85-6P
343574-86-7P
               343574-87-8P
                               343574-88-9P
                                              343574-90-3P
                                                              343574-91-4P
343574-92-5P
               343574-93-6P
                              343574÷94-7P
                                              343574-95-8P
                                                              343574-96-9P
                              343574-99-2P
343574-97-0P
               343574-98-1P
                                              343575-00-8P
                                                              343575-01-9P
                               343575-04-2P
343575-02-0P
               343575-03-1P
                                              343575-05-3P
                                                              343575-06-4P
                                              343575-10-0P
343575-07-5P
               343575-08-6P
                               343575-09-7P
                                                              343575-11-1P
                               343575-15-5P
343575-13-3P
               343575-14-4P
                                              343575-16-6P
                                                              343575-17-7P
                               343575-20-2P
343575-18-8P
               343575-19-9P
                                              343575-21-3P
                                                              343575-22-4P
343575-23-5P
               343575-24-6P
                               343575-25-7P
                                              343575-26-8P
                                                              343575-27-9P
343575-28-0P
               343575-29-1P
                               343575-30-4P
                                              343575-31-5P
                                                              343575-32-6P
                               343575-35-9P
343575-33-7P
               343575-34-8P
                                              343575-36-0P
                                                              343575-37-1P
343575-38-2P
               343575-39-3P
                              343575-40-6P
                                              343575-41-7P
                                                              343575-42-8P
343575-43-9P
               343575-44-0P
                               343575-45-1P
                                              343575-46-2P
               343575-48-4P
                               343575-49-5P
                                              343575-50-8P
343575-47-3P
343575-51-9P
               343575-52-0P
                               343575-53-1P
                                              343575-54-2P
                                                             343575-55-3P
343575-56-4P
               343575-57-5P
                               343575-58-6P
                                              343575-59-7P
                                                             343575-60-0P
               343575-62-2P
343575-61-1P
                              343575-63-3P
                                              343575-64-4P
                                                             343575-65-5P
               343575-67-7P
                               343575-68-8P
                                              343575-69-9P
343575-66-6P
                                                             343575-70-2P
                               343575-73-5P
343575-71-3P
               343575-72-4P
                                              343575-74-6P
                                                              343575-75-7P
               343575-77-9P
                               343575-78-0P
                                              343575-79-1P
343575-76-8P
                                                              343575-80-4P
343575-81-5P
               343575-82-6P
                               343575-83-7P
                                              343575-84-8P
                                                              343575-85-9P
```

建沙罗用金

海岸中四州 田子

ST

ΙT

ΙT

IT

```
343575-89-3P
                    343575-88-2P
                                                    343575-90-6P
                                                                    343575-91-7P
     343575-86-0P
                                                                    343575-97-3P
     343575-93-9P
                    343575-94-0P
                                    343575-95-1P
                                                    343575-96-2P
                                                    343576-02-3P
     343575-99-5P
                    343576-00-1P
                                    343576-01-2P
                                                                    343576-03-4P
                    343576-05-6P
                                    343576-06-7P
                                                    343576-07-8P
                                                                    343576-08-9P
     343576-04-5P
                                    343576-11-4P
                                                    343576-12-5P
                                                                    343576-13-6P
     343576-09-0P
                    343576-10-3P
     343576-14-7P
                    343576-15-8P
    RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (prepn. of 3-(2-aminoethylthio) methyl-4-oxo-4-(3-pyridyl) butanoic acid
        derivs. as neuroprotective agents)
                                                    343576-19-2P
                                                                    343576-20-5P
IT
     343576-16-9P
                    343576-17-0P
                                    343576-18-1P
                                                    343576-24-9P
                                                                    343576-25-0P
     343576-21-6P
                    343576-22-7P
                                    343576-23-8P
     343576-26-1P
                    343576-27-2P
                                    343576-28-3P
                                                    343576-29-4P
                                                                    343576-30-7P
     343576-31-8P
                    343576-32-9P
                                    343576-33-0P
                                                    343576-34-1P
                                                                    343576-35-2P
     343576-36-3P
                    343576-37-4P
                                    343576-38-5P
                                                    343576-39-6P
                                                                    343576-40-9P
     343576-41-0P
                    343576-42-1P
                                    343576-43-2P
                                                    343576-44-3P
                                                                    343576-45-4P
                                                                    343576-50-1P
     343576-46-5P
                    343576-47-6P
                                    343576-48-7P
                                                    343576-49-8P
                    343576-52-3P
                                    343576-53-4P
                                                    343576-54-5P
                                                                    343576-55-6P
     343576-51-2P
                    343576-57-8P
                                    343576-58-9P
                                                    343576-59-0P
                                                                    343576-60-3P
     343576-56-7P
                                    343576-63-6P
                                                                    343576-65-8P
                                                    343576-64-7P
     343576-61-4P
                    343576-62-5P
                                    343576-68-1P
                                                                    343576-70-5P
                    343576-67-0P
                                                    343576-69-2P
     343576-66-9P
                                                                    343576-75-0P
                                                    343576-74-9P
     343576-71-6P
                    343576-72-7P
                                    343576-73-8P
     343576-76-1P
                    343576-77-2P
                                    343576-78-3P
                                                    343576-79-4P
                                                                    343576-80-7P
                                    343576-83-0P
                                                                    343576-85-2P
     343576-81-8P
                    343576-82-9P
                                                    343576-84-1P
                     343576-87-4P
                                    343576-88-5P
                                                    343576-89-6P
                                                                    343576-90-9P
     343576-86-3P
                     343576-92-1P
                                    343576-93-2P
                                                    343576-94-3P
                                                                    343576-95-4P
     343576-91-0P
                                    343576-98-7P
                                                    343576-99-8P
                                                                    343577-00-4P
     343576-96-5P
                     343576-97-6P
                                    343577-03-7P
                                                    343577-04-8P
                                                                    343577-05-9P
     343577-01-5P
                     343577-02-6P
     343577-06-0P
                     343577-07-1P
                                    343577-08-2P
                                                    343577-09-3P
                                                                    343577-10-6P
     343577-11-7P
                                                                    343577-15-1P
                     343577-12-8P
                                    343577-13-9P
                                                    343577-14-0P
                    343577-19-5P
                                    343577-20-8P
                                                    343577-21-9P
                                                                    343577-22-0P
     343577-18-4P
                                    343577-26-4P
                                                    343577-27-5P
                                                                    343577-28-6P
     343577-24-2P
                    343577-25-3P
                                    343577=31-1P
                                                    343577-32-2P
                                                                    343577-33-3P
                    343577-30-0P
     343577-29-7P
                                    343577-36-6P
                                                    343577-37-7P
                                                                    343577-38-8P
                     343577-35-5P
     343577-34-4P
                                                                    343577-43-5P
                     343577-40-2P
                                    343577-41-3P
                                                    343577-42-4P
     343577-39-9P
                                                    343577-47-9P
                                    343577-46-8P
                                                                    343577-48-0P
     343577-44-6P
                     343577-45-7P
     343577-49-1P
                    ·343577-50-4P
                                    343577-51-5P
                                                    343577-52-6P
                                                                    343577-53-7P
     343577-54-8P
                     343577-55-9P
                                    343577-56-0P
                                                    343577-57-1P
                                                                    343577-58-2P
                     343577-60-6P
                                    343577-61-7P
                                                    343577-62-8P
                                                                    343577-63-9P
     343577-59-3P
     343577-64-0P
                     343577-65-1P
                                    '343577-66-2P
                                                    343577-67-3P
                                                                    343577-68-4P
                     343577-70-8P
                                    343577-71-9P
                                                    343577-72-0P
                                                                    343577-73-1P
     343577-69-5P
                                    343577-76-4P
                                                    343577-77-5P
                                                                    343577-78-6P
     343577-74-2P
                     343577-75-3P
                    343577-80-0P
                                    343577-81-1P
                                                    343577-82-2P
                                                                    343577-83-3P
     343577-79-7P
                     343577-85-5P
                                    343577-86-6P
                                                    343577-87-7P
                                                                    343577-88-8P
     343577-84-4P
                                                    343577-93-5P
                     343577-90-2P
                                    343577-92-4P
                                                                    343577-94-6P
     343577-89-9P
                                    343577-97-9P
                    343577-96-8P
                                                    343577-99-1P
                                                                    343578-00-7P
     343577-95-7P
                                    343578-03-0P
                                                                    343578-05-2P
     343578-01-8P
                     343578÷02-9P
                                                    343578-04-1P
                                                                    343578-10-9P
     343578-06-3P
                     343578-07-4P
                                    343578-08-5P
                                                    343578-09-6P
                     343578-12-1P
                                    343578-13-2P
                                                    343578-14-3P
                                                                    343578-15-4P
     343578-11-0P
                                    343578-18-7P
                                                    343578-19-8P
                                                                    343578-20-1P
     343578-16-5P
                     343578-17-6P
                                    343578-23-4P
                                                    343578-24-5P
                                                                    343578-25-6P
                     343578-22-3P
     343578-21-2P
                     343964-43-2P
     343578-31-4P
     RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (prepn. of 3-(2-aminoethylthio) methyl-4-oxo-4-(3-pyridyl) butanoic acid
        derivs. as neuroprotective agents)
  71-23-8, n-Propanol, reactions 86-84-0, 1-Naphthyl isocyanate
                                                                          96-33-3.
                        104-12-1, 4-Chlorophenyl isocyanate
                                                               104-97-2,
     Methyl acrylate
                                     ja Harrit
```

14

121 141

Cyclopentanepropanoyl chloride 110-89-4, Piperidine, reactions 140-88-5, Ethyl acrylate 141-32-2 500-22-1, 3-Pyridinecarboxaldehyde 501-53-1, Benzyl chloroformate 622-58-2, 4-Methylphenyl isocyanate 700-87-8, 2-Methoxyphenyl isocyanate 1016-19-9, 3,4,5-Trimethoxyphenyl 1190-73-4, n-Acetyl cysteamine 1195-45-5, 4-Fluorophenyl 1476-23-9, Allyl isocyanate 2859-67-8, 3-Pyridinepropanol isocyanate isocyanate 3025-95-4 3173-53-3, Cyclohexyl isocyanate 4192-31-8, 4-Oxo-4-(3-pyridyl)butanoic acid 4530-20-5 6068-72-0, 4-Cyanol chloride 7652-46-2 15159-40-7, Morpholine-4-carbonyl chloride 6068-72-0, 4-Cyanobenzoyl 16744-98-2, 2-Fluorophenyl isocyanate 20938-74-3 36643-74-0, Indanol 54132-75-1, 3,5-Dimethylphenyl isocyanate 59587-09-6 67385-09-5 129714-97-2, 3,5-Difluorobenzoyl chloride 220301-12-2 69812-29-9 343578-29-0 343578-30-3 343578-28-9 RL: RCT (Reactant) (prepn. of 3-(2-aminoethylthio)methyl-4-oxo-4-(3-pyridyl)butanoic acid derivs. as neuroprotective agents)

4. 线重新点 44 19 5 11 11

3年:道籍:

59086-27-0P 61192-47-0P 343573-70-6P 343573-71-7P 343573-72-8P

IT 343578-26-7P 343578-27-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of 3-(2-aminoethylthio)methyl-4-oxo-4-(3-pyridyl)butanoic acid derivs. as neuroprotective agents)

IT 343575-47-3P

1 1 RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 3-(2-aminoethylthio) methyl 4-oxo-4-(3-pyridyl) butanoic acid derivs. as neuroprotective agents)
3575-47-3 HCAPLUS
DEX NAME NOT YET ASSIGNED

RN 343575-47-3 HCAPLUS : 1-

CN INDEX NAME NOT YET ASSIGNED

Absolute stereochemistry.

四次第二次第二次 计 ANSWER 2 OF 24 HCAPLUS COPYRIGHT 2001 ACS L27 河洋海峡河北洋

ΑN 2000:592548 HCAPLUS

DN 133:177486

**长和严申报**】。 h Preparation of substituted stilbene compounds with vascular damaging ΤI April 1947 activity

IN Davis, Peter David

PA Angiogene Pharmaceuticals Ltd., UK

SO PCT Int. Appl., 31 pp. CODEN: PIXXD2

DΤ Patent

LA English

IC ICM A61K031-195

ICS A61P035-00; A61P017-00; A61P027-02

130

34-2 (Amino Acids, Peptides, and CC

Proteins)

HARLEY HE

```
FAN.CNT 1
                        KIND DATE
                                               APPLICATION NO.
     PATENT NO.
                               20000824 WO 2000-GB503
     WO 2000048590
                         A1
                                                                   20000215
ΡI
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
              AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
              DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
              CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRAI GB 1999-3403
                         A 19990216
     MARPAT 133:177486
os
     A vascular damaging agent AXB (A = substituted cis-stilbene; X = linker
AB
     bond, atom, or group; B = moiety derived from an inhibitor of the
     formation or action of NO in mammalian systems), is claimed. Thus,
     (Z)-1-[3-(N-.alpha.-tert-butoxycarbonyl-N-.omega.-nitroarginyloxy)-4-
     methoxyphenyl]-2-(3,4,5-trimethoxyphenyl)ethene was stirred with CF3CO2H
     in CH2Cl2 to give (Z)-1-(4-methoxy-3-NG-nitroarginyloxyphenyl)-2-(3,4,5-
     trimethoxyphenyl)ethene. The latter at 50 mg/kg i.p. in mice bearing CaNT
     or SaS tumors gave 95% redn. in vascular vol. and 91-100% tumor necrosis.
     vascular damaging agent substituted stilbene prepn; neovascularization
ST
     disease treatment substituted stilbene; nitroarginyloxystilbene prepn
     tumor vascular damaging activity; nitric oxide synthase inhibitor
     nitroarginyloxystilbene prepn
Angiogenesis
IT
         (neovascularization, treatment of diseases involving
        neovascularization; prepn. of substituted stilbene compds. with
         vascular damaging activity) ( ) 程度 [相]
IT
     Amino acids, preparation
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
                                        11794, 14
         (nitroarginyloxystilbene derivs.; prepn. of substituted stilbene
         compds. with vascular damaging activity)
                                      化多类 经基本债券
ΙT
     Antitumor agents
                           1.7
         (prepn. of substituted stilbene compds. with vascular damaging
         activity)
     125978-95-2, Nitric oxide synthase
IT
     RL: BSU (Biological study, unclassified); MSC (Miscellaneous); BIOL
     (Biological study)
         (inhibitors; prepn. of substituted stilbene compds. with vascular
                                       311 to 1
         damaging activity):
IT
     288585-54-6P 288585-55-7P 288585-56-8P
     288585-57-9P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
     (Preparation); USES (Uses)
         (prepn. of substituted stilbene compds. with vascular damaging
         activity)
                    61240-20-8, 3,4,5-Trimethoxybenzyltriphenylphosphonium
IT
     51298-62-5
                 97315-18-9 117048-59-61 288585-58-0
     bromide
     RL: RCT (Reactant)
         (prepn. of substituted stilbene compds. with vascular damaging
         activity)
                                      Alter in the
     288585-59-1P
                      288585-60-4P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
```

Section cross-reference(s): 1, 25

· I the second

(prepn. of substituted stilbene compds. with vascular damaging activity)

RE.CNT 5

RE

- (1) Ajinomoto Kk; EP 0641767 A 1995 HCAPLUS
- (2) Aston Molecules Ltd; WO 9216486 A 1992 HCAPLUS
- (3) George, R; ANTI-CANCER DRUG DESIGN 1995; V10(4), P299
- (4) Koji, O; JOURNAL OF MEDICINAL CHEMISTRY 1998, V41(16), P3022
- (5) Ohsumi, K; BIOORGANIC & MEDICINAL CHEMISTRY LETTERS 1998, V8(22), P3153 HCAPLUS
- IT 288585-55-7P 288585-56-8P 288585-57-9P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of substituted stilbene compds. with vascular damaging activity)

- RN 288585-55-7 HCAPLUS
- CN L-Ornithine, N5-[imino(nitroamino)methyl]-N2-[[2-methoxy-5-[(1Z)-2-(3,4,5-trimethoxyphenyl)ethenyl]phenoxy]carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

- RN 288585-56-8 HCAPLUS
- CN L-Ornithine, N5-[imino(nitroamino)methyl]-N2-[[2-methoxy-5-[(1Z)-2-(3,4,5-trimethoxyphenyl)ethenyl]phenoxyj[carbonyl]-, (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

- RN 288585-57-9 HCAPLUS
- CN L-Ornithine, N5-[imino(nitroamino)methyl]-N2-[[2-methyl-5-[(1Z)-2-(3,4,5-trimethoxyphenyl)ethenyl]phenoxy]carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

```
ANSWER 3 OF 24 HCAPLUS
                              COPYRIGHT 2001 ACS
L27
                                  The last the
     1999:753019 HCAPLUS:
ΑN
DN
     132:12506
     Preparation of peptides for treating diseases of excessive bone loss or
ΤI
     cartilage or matrix degradation as cysteine protease inhibitors
     Bondinell, William Edward; Desjarlais, Renee Louise; Veber, Daniel Frank;
IN
     Yamashita, Dennis Shinji
PA
     Smithkline Beecham Corporation, USA
SO
     PCT Int. Appl., 128 pp.
     CODEN: PIXXD2
                                  CALCARD N
DT
     Patent
LА
     English
     ICM A61K
IC
CC
     34-3 (Amino Acids, Peptides, and
     Section cross-reference(s): 1, 7, 63
FAN.CNT 1
     PATENT NO.
                      KIND'
                            DATE
                                          APPLICATION NO.
                                                           DATE
                                       WO 1999-US11266
                                                           19990520
PΙ
     WO 9959526
                      A2
                            19991125
                      A3
     WO 9959526
                           20000120:
            AE, AL, AU, BA, BB, BG, BR, CA, CN, CZ, EE, GE, GH, GM, HR, HU,
            ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MG, MK, MN, MX,
            NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA,
            AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                      A2 20010117 EP 1999-924421
     EP 1067894
         R: BE, CH,
                    DE, ES, FR, GB, IT, LI, NL
```

a free

Fil

"就走"。 特

144.7

os

GΙ

$$X = R^4 \longrightarrow NH \longrightarrow R^3 \longrightarrow R^3$$

The present invention provides peptides bis-aminomethyl carbonyl protease AB inhibitors I (R1, R2 = alkyl; P = X, Y; R3 selected from the group consisting of: CH2CH(CH3)2, CH2CH2CH3, CH2CH=CH2, or CH2Ph; R4 is selected from the group consisting of alkyl; N-piperazine; Ntetrahydroisoquinoline; substituted alkyl, Ph, benzofuran, benzothiazole; quinoline; naphthyl; and benzoxazole; R5 = Ph and Ph substituted with alkyl, N-piperidine, benzofuran; pyridine; Q = arylacyl) and pharmaceutically acceptable salts, hydrates and solvates thereof which inhibit proteases, including cathepsin K, pharmaceutical compns. of such compds., and methods for treating diseases of excessive bone loss or cartilage or matrix degrdn., including osteoporosis; gingival disease including gingivitis and periodontitis; arthritis, more specifically, osteoarthritis and rheumatoid arthritis; Paget's disease; hypercalcemia of malignancy; and metabolic bone disease, comprising inhibiting said bone loss or excessive cartilage or matrix degrdn. by administering to a patient in need thereof a compdate of the present invention. Thus,  $(S)-3N-(N-(thianaphthenyl-2-carbonyl)-leucinyl)-amino-1N-(3-{2-(1-oxo)-1})$ pyridyl}phenylacetyl)-amino-butan-2-one was prepd. for treating diseases of excessive bone loss or cartilage or matrix degrdn. as cysteine protease inhibitor. Detn. of cathepsin K proteolytic catalytic activity of these compds. are reported.

ST osteoarthritis osteoporosis rheumatoid arthritis peptide prepn; bone loss cartilage degrdn peptide prepn protease inhibitor

IT Cartilage

(degeneration; prepn. of peptides for treating diseases of excessive bone loss or cartilage or matrix degrdn. as cysteine protease inhibitors)

ુવા મુનિયુ તાલું ક

John the Har

Charmetta, or II

IT Bone

(demineralization; prepn. of peptides for treating diseases of excessive bone loss or cartilage or matrix degrdn. as cysteine protease inhibitors)

IT Bone, disease
Osteoarthritis
Osteoporosis
Rheumatoid arthritis

(prepn. of peptides for treating diseases of excessive bone loss or cartilage or matrix degrdn. as cysteine protease inhibitors)

im the sign of

Mi tomalor.

IT Peptides, preparation

RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of peptides for treating diseases of excessive bone loss or cartilage or matrix degrdn. as cysteine protease inhibitors)

IT 251458-51-2P

RL: BAC (Biological activity or effector, except adverse); RCT (Reactant);

Little kare o in the section of the

Till in Hi

```
SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
         (prepn. of peptides for treating diseases of excessive bone loss or
        cartilage or matrix degrdn as cysteine protease inhibitors) 119-79-5P 247119-80-8P 247119-82-0P 247119-83-1P 251
                                                                       251457-08-6P
ΙT
     247119-79-5P
                     251457-10-0P
                                     251457-14-4P
                                                       251457-15-5P
                                                                       251457-16-6P
     251457-09-7P
     251457-17-7P
                     251457-18-8P
                                      251457-19-9P
                                                       251457-21-3P
                                                                       251457-23-5P
                                     251457-28-0P
                                                       251457-30-4P
                                                                       251457-32-6P
     251457-24-6P
                     251457-27-9P
                                      251457-36-0P 251457-37-1P
                                                                       251457-38-2P
                     251457-35-9P
     251457-34-8P
                     251457-40-6P
                                      251457-41-7P
                                                       251457-42-8P
                                                                       251457-43-9P
     251457-39-3P
                                      251457-46-2P
                                                       251457-47-3P
                                                                       251457-48-4P
                     251457-45-1P
     251457-44-0P
                                      251457-51-9P
                                                       251457-52-0P
                                                                       251457-53-1P
     251457-49-5P
                     251457-50-8P
                                     251457-56-4P
                                                       251457-60-0P
                                                                       251457-62-2P
     251457-54-2P
                     251457-55-3P
                     251457-65-5P
                                      251457-66-6P
                                                       251457-68-8P
                                                                       251457-69-9P
     251457-64-4P
                                      251457-75-7P 251457-76-8P
     251457-71-3P
                     251457-74-6P
                                                                       251457-77-9P
     251457-78-0P
                     251457-79-1P
                                      251457-80-4P
                                                       251457-82-6P
                                                                       251457-84-8P
                     251457-87-1P
                                      251457-89-3P
                                                       251457-90-6P
                                                                       251457-91-7P
     251457-85-9P
     251457-92-8P
                     251457-93-9P
                                      251457-94-0P 251457-95-1P
                                                                       251457-96-2P
                                                                       251458-02-3P
     251457-97-3P
                     251457-98-4P
                                      251457-99-5P
                                                       251458-01-2P
                                      251458-06-7P
                                                                       251458-09-0P
                     251458-05-6P
                                                       251458-08-9P
     251458-04-5P
                                      251458-12-5P
                                                                       251458-17-0P
     251458-10-3P
                     251458-11-4P
                                                       251458-13-6P
                                     251458=20-5P 251458-24-9P
                                                                       251458-29-4P
                     251458÷19-2P
     251458-18-1P
                     251458 132-9P
                                                       251458-35-2P
                                                                       251458-37-4P
     251458-31-8P
                                     251458-33-0P
                                      251458-40-9P
                     251458-41-0P
                                                                       251458-44-3P
                                      251458-52-3P
                                                                       251458-55-6P
     251458-45-4P
                     251458-50-1P
                                                       251458-53-4P
                     251458-59-0P
                                      251458-61-4P
                                                       251458-62-5P
                                                                       251458-63-6P
     251458-57-8P
                     251458-65-8P
                                                       251458-67-0P
                                      251458-66-9P
                                                                       251458-68-1P
     251458-64-7P
                                      251458-71-6P
                     251458-70-5P
                                                      251458-78-3P
                                                                       251458-79-4P
     251458-69-2P
                                       11:1
     251458-80-7P
     RL: BAC (Biological activity of effector) except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
     (Preparation); USES (Uses)
         (prepn. of peptides for treating diseases of excessive bone loss or
         cartilage or matrix degrdn. (as cysteine protease inhibitors)
     37353-41-6, Cysteine protease RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
ΙT
         (prepn. of peptides for treating diseases of excessive bone loss or
        cartilage or matrix degrdn. as cysteine protease inhibitors)
     50-43-1, 2,4,6-Trichlorobenzoic acid $51-44-5, 3,4-Dichlorobenzoic acid
     65-85-0, Benzoic acid, reactions: 75-08-1, Ethanethiol 79-09-4,
                                    86-59-9, 8-Quinoline-carboxylic acid
     Propionic acid, reactions
     88-13-1, Thiophene-3-carboxylic acid 91-21-4, 1,2,3,4-
     Tetrahydroisoquinoline 92-92-2; Biphenyl-4-carboxylic acid"
     3,4-Dimethoxybenzoic acid 93-09-4, 2-Naphthoicacid
                                                                   93-10-7,
                                      98-09-9, Benzenesulfonyl chloride
     2-Quinolinylcarboxylic acid
     Pyrazine-2-carboxylic acid 99-96-7, 4-Hydroxybenzoic acid, reactions 100-00-5, 1-Chloro-4-nitro-benzene 100-09-4, 4-Methoxybenzoic acid" 109-01-3 109-52-4, Valeric acid, reactions 142-25-6 349-88-2, 4-Fluorophenylsulfonyl chloride 371-40-4, 4-Fluoro-aniline 402-54-0
     455-24-3, 4-(Trifluoromethyl)benzoic acid 456-22-4, 4-Fluoro-benzoic
             496-41-3, Benzofuran-2-carboxylic acid 501-52-0,
     acid
     3-Phenylpropionic acid 504-29-0, 2-Amino-pyridine
                                                                 536-69-6,
     5-Butylpyridine-2-carboxylic acid 585-76-2, 3-Bromo-benzoic acid 611-95-0, 4-(Benzoyl)benzoic acid 616-29-5, 1,3-Diamino-propan-2-ol
     619-45-4, 4-Methoxycarbonyl-aniline: 620-86-0 622-40-2,
     4-(2-\text{Hydroxyethyl}) morpholine 636 \div 82-8, 1-\text{Cyclohexene}-1-\text{carboxylic} acid
                 661-69-8, Hexamethylditin 828-51-3, Adamantane-1-carboxylic
     646-07-1
             883-21-6, 1-Methoxy-2-naphthoicacid 883-62-5,
     3-Methoxy-2-naphthoic acid 1457-59-6; 5-Methylimidazolyl-4-carboxylic
             1477-50-5, Indole-2-carboxylic acid 1486-51-7 1623-92-3,
```

Acres 18 19

特性特殊 化

```
4-Phenoxybenzenesulfonyl chloride 1670-82-2, Indole-6-carboxylic acid
     1723-27-9, Thieno[3,2-b]thiophene-2-carboxylic acid 1878-67-7, 3-Bromo phenyl acetic acid 2008-75-5, 2-(Piperidin-1-yl)ethyl chloride
     hydrochloride"
                                    2991-42-6, 4-Trifluoromethylbenzenesulfonyl
                        2666-93-5
                 3328-70-9, 5-Formylsalicylaldehyde 3510-66-5,
     chloride
     2-Bromo-5-methylpyridine 3622-35-3, 6-Benzothiazolecarboxylic acid
     3647-69-6, 2-(4-Morpholinyl)ethyl chloride hydrochloride 4457-32-3,
     4-Nitrobenzyl chloroformate
                                     4467-07+6, 3-(2-Pyridyl) benzoic acid
     4926-28-7, 2-Bromo-4-methylpyridine 5315-25-3, 2-Bromo-6-methylpyridine
                  6314-28-9, Benzo[b]thiophene-2-carboxylic acid
                                                                         6315-89-5,
     3,4-Dimethoxy-aniline
                               6480-68-8, 3-Quinolinecarboxylic acid
     Methyl 3-hydroxy-4-methoxy-benzoate 6973-60-0
                                                            7250-53-5,
     5-Quinoline-carboxylic acid 10147-37-2, Isopropyl sulfonyl chloride 10349-57-2, 6-Quinolinecarboxylic acid 15112-41-1, 5-Benzoxazolecarboxylic acid 16309-45-8 16419-60-6 19438-10-9, Me
                                                                 19438-10-9, Methyl
                           20029-52-1, 4-Cyclohexylbenzoic acid
     3-hydroxybenzoate
     31462-59-6, Pyrimidine-4-carboxylic acid 32315-10-9, Triphosgene
                  50551-56-9
     40274-67-7
                                50793-29-8, 4-(4-Cyanophenoxy) benzoic acid
     57260-71-6, 1-(tert-Butoxycarbonyl)piperazine 65007-00-3
                                                                         66715-65-9,
                                     68947-43-3, N-Methylpiperidine-4-carboxylic
     2-Pyridinesulfonyl chloride
             73579-08-5
                           78161-82-7, 4-(4-(Trifluoromethyl)phenoxy)benzoic acid
     81432-12-4 90433-20-8 92198-45-3 9 98327-87-8, 2,2'-
Bis (diphenylphosphino)-1,1'-binaphthyl 98437-24-2, 2-Benzofuran boronic
                           150529-73-0, Methyl 3-bromophenylacetate
             144059-86-9
                     154235-77-5, Benzoxazole-6-carboxylic acid
175201-51-1 190661-70-2 210304-58-8 2
     150798-78-0
                                                                       154739-53-4
     175137-58-3
                                                                   215947-98-1
                                    190661-70-2

251457-20-2

251457-33-7

251458-15-8

251458-26-1

251458-30-

251458-30-

251458-30-
     228419-12-3
                     248923-01-5
     251457-88-2
                     251458-03-4
                                                                   251458-30-7
                     251458+38-5
     251458-34-1
                                     بي ڏائي ج
     RL: RCT (Reactant)
         (prepn. of peptides for treating diseases of excessive bone loss or
         cartilage or matrix degrdn. as cysteine protease inhibitors)
ΙT
     23948-77-8P, [1,1'-Biphenyl]-3-acetic acid 51061-68-8P
                                                                      69038-74-0P
                                  203503 06 4P 215309-01-6P
P 224643-30-5P 227178-31-6P
                     99370-68-0P
     75852-28-7P
                                                                     216316-23-3P
                     216316#27-7P
     216316-24-4P
                                                                        227178-32-7P
     229630-60-8P
                     247119-96-6P
                                      247119-99-9P : 247122-09-4P
                                                                        250726-45-5P
                                      251457-05-3P 251457-06-4P 251457-13-3P 251457-22-4P
     250726-46-6P
                     250726-47-7P
                                                                        251457-07-5P
     251457-11-1P
                     251457-12-2P
                                                                        251457-25-7P
     251457-26-8P
                     251457-29-1P
                                      251457-31-5P 251457-57-5P
                                                                        251457-58-6P
                     251457-61-1P
                                      251457-63-3P · 251457-67-7P
     251457-59-7P
                                                                       251457-70-2P
     251457-73-5P
                     251458-00-1P
                                      251458-07-8P
                                                       251458-14-7P
                                                                       251458-16-9P
     251458-21-6P
                     251458-22-7P
                                      251458-23-8P
                                                       251458-25-0P
                                                                       251458-27-2P
                                                       251458-47-6P
                     251458-36-3P
                                      251458-46-5P
                                                                        251458-48-7P
     251458-28-3P
                                      251458-74-9P
     251458-49-8P
                     251458-72-7P
                                                       251458-75-0P
                                                                       251458-76-1P
     251458-77-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. of peptides for treating diseases of excessive bone loss or
        cartilage or matrix degrdn. as cysteine protease inhibitors)
IΤ
     251457-72-4
                                       RL: RCT (Reactant)
         (prepn. of peptides for treating diseases of excessive bone loss or
        cartilage or matrix degrdn. as cysteine protease inhibitors)
                                       随即 排 一种 建二十二
RN
     251457-72-4 HCAPLUS
CN
     L-Leucine, N-[(4-nitrophenoxy)carbonyl]-, 1,1-dimethylethyl ester (9CI)
                                     (CA INDEX NAME)
```

医维维氏硬膜炎 珠色点

明確認的

Absolute stereochemistry.

L27 ANSWER 4 OF 24 HCAPLUS COPYRIGHT 2001 ACS

AN 1998:749736 HCAPLUS

DN 130:110572

TI Self-Immolative Nitrogen Mustard Prodrugs for Suicide Gene Therapy
Niculescu-Duvaz, Dan; Niculescu-Duvaz, Ion; Friedlos, Frank; Martin,
Janet; Spooner, Robert; Davies, Lawrence; Marais, Richard; Springer,
Caroline J.

CS UK

SO J. Med. Chem. (1998), 41(26), 5297-5309 CODEN: JMCMAR; ISSN: 0022-2623

PB American Chemical Society

DT Journal

LA English

CC 34-2 (Amino Acids, Peptides, and Proteins)
Section cross-reference(s): 1, 2

GΙ

Four new potential self-immolative prodrugs derived from phenol and aniline nitrogen mustards, four model compds. derived from their corresponding fluoroethyl analogs and two new self-immolative linkers were designed and synthesized for use in the suicide gene therapy termed GDEPT (gene-directed enzyme prodrug therapy). The self-immolative prodrugs were designed to be activated by the enzyme carboxypeptidase G2 (CPG2) releasing an active drug by a 1,6-elimination mechanism via an unstable intermediate. Thus, glutamate derivs. I (Y = O, NH; Y = O, NH; X = Cl)

were synthesized. They are bifunctional alkylating agents in which the activating effects of the phenolic hydroxyl or amino functions are masked through an oxycarbonyl or a carbamoyl bond to a benzylic spacer which is itself linked to a glutamic acid by an oxycarbonyl or a carbamoyl bond. The corresponding fluoroethyl compds. I (X = F) were also synthesized. The rationale was to obtain model compds. with greatly reduced alkylating abilities that would be much less reactive with nucleophiles compared to the corresponding chloroethyl derivs. This enabled studies of these model compds. as substrates for CPG2; without incurring the rapid and complicated decompn. pathways of the chloroethyl derivs. The prodrugs were designed to be activated to their corresponding phenol and aniline nitrogen mustard drugs by CPG2 for use in GDEPT. The synthesis of the analogous novel parent drugs II is also described. A colorectal cell line was engineered to express CPG2 tethered to the outer cell surface. The phenylenediamine compds. were found to behave as prodrugs, yielding IC50 prodrug/IC50 drug ratios between 20- and 33-fold for I (X = Cl, Y = NH; Z = 0, NH) and differentials of 12-14-fold between CPG2-expressing and control LacZ-expressing clones. The drugs released are up to 70-fold more potent than benzoic acid III that results from the glutamate prodrug (CMDA) which has been used previously for GDEPT. These data demonstrate the viability of this strategy and indicate that self-immolative prodrugs can be synthesized to release potent mustard drugs selectively by cells expressing CPG2 tethered to the cell surface in GDEPT. glutamate nitrogen mustard prodrug preph suicide gene therapy; immolative nitrogen mustard prodrug suicide gene therapy; carboxypeptidase G2 cleavage glutamate nitrogen mustard prodrug prepn; anticancer agent glutamate nitrogen mustard prodrug preph; cytotoxicity glutamate nitrogen mustard prodrug prepn motherapy
(gene-directed enzyme prodrug; prepn. of self-immolative nitrogen Chemotherapy mustard prodrugs for suicide gene therapy) · 🕴 - -Antitumor agents . f., usef a t Cytotoxicity 护制技术 Gene therapy Prodrugs (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene therapy) tradition tops of it 122665-70-7 RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study) (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene 1204-69-9, 4-[Bis(2-chloroethyl)amino]phenol : 2067-58-5 RL: BAC (Biological activity of effector, except adverse); RCT (Reactant); BIOL (Biological study) (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene 219592-00-4P therapy) 219591-87-4P RL: BAC (Biological activity or effector, except adverse); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene 122665-73-0DP, N-4-[(2-Chloroethyl)(2-mesyloxyethyl)amino]benzoyl-Lglutamic acid, dichloro and difluoro carbonate and carbamate analogs 180839-05-8P 219591-76-1P 180838-98-6P 180839-01-4P 180839-03-6P 219591-94-3P 219591-82-9P 219591-86-3P 219591-92-1P RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene

1-1-1-14

therapy)

ST

IT

IT

ΙT

IT

IT

IT

```
9074-87-7, Carboxypeptidase G2
IT
    RL: BPR (Biological process); BTOL (Biological study); PROC (Process)
        (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene
       therapy)
                   219591-79-4P
    219591-77-2P
IT
    RL: BYP (Byproduct); PREP (Preparation)
        (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene
                                  1 713 35.
       therapy)
                                  AND BUTT
    219591-73-8P
ΙT
    RL: BYP (Byproduct); RCT (Reactant); PREP (Preparation)
        (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene
        therapy)
    123-08-0, 4-Hydroxybenzaldehyde 619-73-8, 4-Nitrobenzyl alcohol
ΙT
                 20845-16-3
                             32677-01-3, Di-tert-butyl L-glutamate
    18226-17-0
                    101582-69-8
    hydrochloride
    RL: RCT (Reactant)
        (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene
        therapy)
                                 57529-05-2P
ΙT
    18483-99-3P
                   18484-05-4P
                                              113068-95-4P
                                                              161803-03-8P
                                   161803-06-1P 180838-97-5P
    161803-04-9P
                   161803-05-0P
                                                                 180839-00-3P
                    180839-04-7P
                                   180839-06-9P
                                                  180839-11-6P
                                                                 180839-13-8P
    180839-02-5P
                                  180839-16-1P
    180839-14-9P
                   180839+15-0P
                                                  180839-18-3P
                                                                 180839-19-4P
    180839-20-7P 180839-21-8P
                                180841-62+7P_{\odot}, 219591-81-8P_{\odot}
                   219591-85-2P
    219591-83-0P
                                   219591-91-0P 219591-93-2P
                                                                 219591-96-5P
                   219591-98-7P
                                   219592-01-5P
                                                  219592-02-6P
    219591-97-6P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene
                                   111
                        · . .
       therapy)
                   219591-89-6P
    219591-75-0P
IT
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene
                                   therapy)
RE.CNT
                                      RE
(1) Alexander, R; Tetrahedron Lett 1991, V32, P3269 HCAPLUS
(2) Bridgewater, G; Eur J. Cancer 1995, V31A, P2362
(3) Carl, P; J Med Chem 1981, V24, P479 HCAPLUS
(4) Corey, E; J Am Chem Soc 1972, V94, P6190 HCAPLUS
(5) Dowell, R; J Med Chem 1996, V39, P1100 HCAPLUS
(6) Dubowchik, G; Tetrahedron Lett 1997, V38, P5261 HCAPLUS
(7) Friedlos, F; J Med Chem 1997, V40, P1270 HCAPLUS
(8) Greene, T; Protective groups in organic chemistry 1991
(9) Hanessian, S; Can J Chem 1975, V53, P2975 HCAPLUS
(10) Hollitzer, O; Angew Chem Int Ed Engl 1976, V5, P444
(11) Huber, B; Proc Natl Acad Sci USSA 1991, V88, P8039 HCAPLUS
(12) Jarman, M; Chem Br 1989, V25, P51 HCAPLUS
(13) Kruse, C; J Org Chem 1979, V44, P1847 HCAPLUS
(14) Marais, R; Cancer Res 1996, V56, P4735 HCAPLUS
(15) Marais, R; Nature Biotechnol 1997, V15, P1373 HCAPLUS
(16) Minton, N; J Bacteriol 1983, V156, P1222 HCAPLUS
(17) Niculescu-Duvaz, I; Bioconjugate Chem 1998, V9, P4 HCAPLUS
(18) Olofson, R; Tetrahedron Lett 1980, V21, P819 HCAPLUS
(19) Palmer, B; J Med Chem 1990, V33, P112 HCAPLUS
(20) Perrone, E; EP 0236880 A2 1987 HCAPLUS
(21) Ross, W; Biological alkylating agents 1962
(22) Schmidt, F; Bioorg Med Chem Lett 1997, V7, P1071 HCAPLUS
(23) Schmidt, H; Chem Ber 1979, V112, P727 HCAPLUS
(24) Sherwood, R; Eur J Biochem 1985, V24, P447
(25) Springer, C; J Med Chem 1990, V33, P677 HCAPLUS
(26) Springer, C; J Med Chem 1994, V37, P2361 HCAPLUS
```

356年前往上15日1日 1

· 是有人的 (1)

前身是"严"的方式

. .

- (27) Springer, C; J Med Chem 1995, V38, P5051 HCAPLUS
- (28) Wakselman, M; N J Chim 1983, V7; P439 HCAPLUS
- (29) Zhang, W; Adv Pharmacol (San Diego) 1995; V6, P289

## IT 180839-20-7P 180839-21-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of self-immolative nitrogen mustard prodrugs for suicide gene therapy)

RN 180839-20-7 HCAPLUS

CN L-Glutamic acid, N-[[4-[[(4-nitrophenoxy)carbonyl]oxy]methyl]phenoxy]carbonyl]-, bis(1,1-dimethylethyl) ester (9CI) (CA INDEX NAME)

#### Absolute stereochemistry.

RN 180839-21-8 HCAPLUS

CN L-Glutamic acid, N-[[4-[[[(4-nitrophenoxy)carbonyl]oxy]methyl]phenoxy]carbonyl]-, di-2-propenyl ester (9CI) (CA INDEX NAME)

#### Absolute stereochemistry.

PAGE 1-B

CH<sub>2</sub>

L27 ANSWER 5 OF 24 HCAPLUS COPYRIGHT 2001 ACS

AN 1998:147326 HCAPLUS

DN 128:205147

```
TΙ
         Preparation of non-peptide bombesin receptor antagonists
         Horwell, David Christopher; Pritchard, Martyn Clive
IN
         Warner-Lambert Company, USA; Horwell, David Christopher; Pritchard, Martyn
PA
         Clive
         PCT Int. Appl., 112 pp.
SO
         CODEN: PIXXD2
         Patent
DT
         English
LΑ
                                                                 1.0
         ICM C07D401-12
IC
                 A61K031-395; C07D209-20; C07D213-89; C07D213-40; C07D213-56;
                  C07D405-12; C07D417-12
CC
         34-3 (Amino Acids, Peptides, and ...
         Section cross-reference(s): 1 ***
FAN.CNT 1
         PATENT NO.
                                       KIND DATE
                                        A1 19980226 WO 1997-US13871 19970806
PΙ
         WO 9807718
                W: AL, AU, BA, BB, BG, BR, CA, CN, CZ, EE, GE, GH, HU, IL, IS, JP, KR, LC, LK, LR, LT, LV, MG, MK, MN, MX, NO, NZ, PL, RO, SG, SI,
                       SK, SL, TR, TT, UA, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD,
                       RU, TJ,
                                            100
                                     TM
                RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NE, PT, SE, BF, BJ, CF, CG, CI, CM, GA,
                       A 19990817
                                                                            BR 1997-11342
                                                                                                            19970222
         BR 9711342
                                                  19980306
                                         A1 .
                                                                            AU 1997-41466
                                                                                                            19970806
         AU 9741466
         AU 733226
                                        B2 20010510
                                        A1 19990609 EP 1997-939359
                                                                                                            19970806
         EP 920424
                    AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                       IE, FI
                                         T2 20010123
                                                                             JP 1998-510779
                                                                                                            19970806
         JP 2001500850
                                      B1 20010227
                                                                         US 1999-230933
                                                                                                            19990203
         US 6194437
                                                                       NO 1999-788
                                        A 19990219
                                                                                                            19990219
         NO 9900788
                                         P 🚯 19960822 👯
PRAI US 1996-24323
                                        W 4 19970806
         WO 1997-US13871
                                                                             N. H. Signer
         MARPAT 128:205147
OS
         Compds. Ar(CR1R8)0-1(CH2)0-1NR4CONR5CR7(CH2Ar1)CONR6(CH2)0-3(CR2R9)0-
AB
         1(CH2)0-2R3 [Ar = Ph, (un)substituted pyridyl; R1, R2 = H, alkyl,
         cycloalkyl; R8, R9 = H or forms a ring with R1 or R2, resp; Ar1 = Ar or
         pyridyl-N-oxide, indolyl, pyridyl; imidazole; R4, R5, R6, R7 = H, Me; R3 =
         Ar or H, OH, Me2N, N-methylpyrrole, etc. or their pharmaceutically
         acceptable salts were prepd. as bombesin receptor antagonists. Thus,
         2-[3-(2,6-diisopropylphenyl)ureldo]-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methyl-N-(1-3-(1+indol-3-yl)-2-methy
         pyridin-2-ylcyclohexylmethyl) propionamide was prepd. by condensation of
         .alpha.-methyl-L-tryptophan with 2,6-diisopropylphenyl isocyanate,
         followed by amidation with 1-pyridin-2-ylcyclohexylmethylamine. Affinity
         binding data (IC50 values) for the product were detd. to be 5 and <10 nM
         for the NMB and GRP receptors, resp.
         pseudopeptide prepn bombesin receptor antagonist
ST
                                                            国。[13] [4] [4]
IT
         Bombesin receptors
         RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
               (prepn. of non-peptide bombesin receptor antagonists)
IT
         Peptides, preparation
         RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
         preparation); THU (Therapeutic use); BIOL (Biological study); PREP
         (Preparation); USES (Uses)
               (pseudopeptides; prepn. of non-peptide bombesin receptor antagonists) 066-87-5P 204067-04-9P
IT
         RL: BAC (Biological activity of effector, except adverse); RCT (Reactant);
                                                               In the state of the first
```

自由學事。自由學生 人名英

· 有种性性的现在分词

```
SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
        (prepn. of non-peptide bombesin receptor antagonists)
                    185215-75-2P
                                   204066-70-6P
                                                  204066-71-7P
                                                                  204066-72-8P
IT
     142627-75-6P
                                   204066-75-1P 204066-76-2P
                                                                  204066-77-3P
     204066-73-9P
                    204066-74-0P
                                   204066-80-8P 204066-81-9P
     204066-78-4P
                    204066-79-5P
                                                                  204066-82-0P
     204066-83-1P
                    204066-84-2P
                                   204066-85-3P
                                                 204066-86-4P
                                                                  204066-89-7P
                    204066-93-3P
                                   204066-95-5P
                                                  204066-97-7P
                                                                  204066-99-9P
     204066-91-1P
                                   204067-03-8P
                    204067-02-7P
                                                                  204067-06-1P
     204067-01-6P
                                                 1204067-05-0P
                    204067-40-3P
     204067-38-9P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
    preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); USES (Uses)
        (prepn. of non-peptide bombesin receptor antagonists)
IT
     204067-30-1P
     RL: BYP (Byproduct); PREP (Preparation)
        (prepn. of non-peptide bombesin receptor antagonists)
IT
     54-12-6, Tryptophan 62-53-3, Benzenamine, reactions
                               100-28-7, p-Nitrophenyl isocyanate
                                                                     111-24-0,
     L-Tryptophan, reactions
                                                                   487-59-2
     1,5-Dibromopentane 153-91-3, Tryptophan, .alpha. methyl-
                 1206-13-9
                                        2739-97-1, 2-Pyridylacetonitrile
                             1548-13-6
     1121-78-4
     3218-02-8, Cyclohexanemethanamine 1 4000-72-0 4442-85-7,
     2-Cyclohexylethylamine 5766-79-0 7693-46-1, p-Nitrophenyl
     chloroformate
                     16709-25-4, alpha.-Methyl-L-tryptophan 17540-18-0
     18502-05-1, 4-Imidazoleacetonitrile 23357-52-0 24544-04-5, 2,6-Diisopropylaniline 25756-29-0, n-Methylcyclohexylmethylamine
     28178-42-9, 2,6-Diisopropylphenyl isocyanate
                                                    30806-83-8
                  50528-53-5;
                               56452-52-9 61341-86-4, s 1-Aminoindan
     40465-45-0
                               103110-98-1 114524-80-0 114779-79-2
                  81428-13-9
     70258-01-4
     135207-25-9
                   142854-50-0
                                 164323-86-8 4 204067-18-5
                                                              204067-20-9
     204067-24-3
                   204067-28-7
    RL: RCT (Reactant)
        (prepn. of non-peptide bombesin receptor antagonists)
ΙT
                                13458-33-8P
                                             35392-66-6P
                                                             37982-29-9P
     5815-73-6P
                  13139-14-5P
                                 75342+32-4P 1 75342-33-5P
                                                             127978-70-5P
     55270-47-8P
                   64464-46-6P
                    135627-42-8P
                                   142946-11-0P
                                                  187610-49-7P
                                                                  187610-50-0P
     135627-41-7P
     190333-61-0P
                    204067-07-2P
                                   204067-08-3P
                                                  204067-09-4P
                                                                  204067-10-7P
                                   204067-13-0P
     204067-11-8P
                    204067-12-9P
                                                  204067-14-1P
                                                                  204067-15-2P
                                   204067-19-6P 204067-21-0P
     204067-16-3P
                    204067-17-4P
                                                                  '204067-22-1P
     204067-23-2P
                    204067-25-4P
                                   204067-26-5P 204067-27-6P
     204067-29-8P
                    204067-31-2P
                                   204067-32-3P
                                                  204067-33-4P
                                                                  204067-34-5P
                                   204067-37-8P
     204067-35-6P
                    204067-36-7P
                                                  204067-39-0P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of non-peptide bombesin receptor antagonists)
IT
     204067-27-6P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of non-peptide bombesin receptor antagonists)
RN \cdot
     204067-27-6 HCAPLUS!
    Tryptophan, .alpha.-methyl-N-[(4-nitrophenoxy)carbonyl]-, methyl ester
CN
                                   (9CI) (CA INDEX NAME)
                 0
```

手超到微微的 辩

Marian da de la companya da de la companya da de la companya de la

resp. D:

间槽

- 1998:17976 HCAPLUS AN thread in Jeph 12 to DN 128:61798 Preparation of epoxide peptidomimetics as irreversible HIV protease ΤI inhibitors Yoon, Heungsik; Choy, Nakyen; Kim, Sung Chun; Choi, Ho Il; Son, Young IN Chan; Park, Chi Hyo; Moon, Kwang-yul; Jung, Wonhee; Kim, Chung Ryeol; Lee, Chang Sun; Koh, Jong Sung; Kim, Sang Soo PA LG Chemical Ltd., S. Korea U.S., 50 pp. Cont.-in-part of U.S. Ser. No. 341,352, abandoned.
- so CODEN: USXXAM

ANSWER 6 OF 24 HCAPLUS COPYRIGHT 2001 ACS

DT Patent English LΑ IC ICM A61K031-44

L27

- ICS A61K031-47 NCL 514314000
- CC 34-3 (Amino Acids, Peptides, and · i . Section cross-reference(s): 1, 63

#### PATENT NO. KIND DATE A 19971209 US 1995-473877 A 19961224 US 1993-159382 B1 19971205 KR 1994-13423 A 19980630 US 1995-572402 A 19980428 US 1996-667888 US 5696134 19950607 PIUS 5587388 19931130 KR 125117 19940615 19951214 US 5773468 19960620 US 5744621 A 19980609 US:1996-667133 19960620 US 5763631 A2 / 19931130 表示性证明 。 PRAI US 1993-159382 A 19940615 KR 1994-13423 B2 19941117 Fair Fair US 1994-341352 A 19921202 KR 1992-23088 A 19921202 A 19930614 KR 1992-23089 KR 1993-10811 A 19931014 KR 1993-21298 19931014 KR 1993-21299 Α KR 1993-21300 A 19931014 US 1995-473877 A2 | 19950607 19951026 KR 1995-37292 os MARPAT 128:61798

## STRUCTURE DIAGRAM TOO LÄRGE FOR DISPLÄY 🕀 AVÄILABLE VIA OFFLINE PRINT \*

Novel cis-epoxide compds. I [R1; R2 = independently H, alkyl; R3 = aryl or AB alkyl (un)substituted; with arom (1/C3-8) cycloalkyl; R4 = H, C1-4 alkyl; n = 0-2; X = CO, COCO, S(0), SO2,  $(S_3)(Y) = (0, 1)$  CH2, NH, NMe; m = 0, 1; R5 = heterocycle; straight, branched, or cyclic C1-8 alkyl; alkyl substituted with heterocycle or cycloalkyl; straight; branched, or cyclic C1-8 alkoxy; aryl-substituted alkoxy; NR6R7; R6 = straight or branched C1-8 alkyl, cycloalkyl, alkyl substituted with cycloalkyl; R7 = H, alkyl; Z = O, NH, NMe; R8, R9 = independently alkyl (un) substituted with arom. hydrocarbon or cycloalkyl; C3-8 cycloalkyl; arom; [ are useful for treating or preventing diseases caused by HIV infection. The novel HIV protease inhibitors I have specific structures to form stable bonding with the enzyme active site, which entails a highly enhanced irreversible

GΙ

```
inhibition against HIV protease. Thus deprotection and peptide coupling
of olefin II (prepd.:in: 4 steps: from protected L-phenylalaninal and
(S)-2-amino-3-methyl-1-phenylbutane) with penicillamine-derived sulfone
III (prepd. in 3 steps from L-penicillamine), followed by epoxidn. with
mCPBA gave title epoxide deriv: TV: TV showed irreversible inactivation
of HIV-1 protease, with a stoichiometric ratio of inhibitor to enzyme of
1:1. IV also showed antiviral activity against HIV-1 with IC50 = 1 nM.
epoxide peptidomimetic prepn HIV protease inhibitor; virucide HIV epoxide
peptidomimetic prepn; AIDS treatment epoxide peptidomimetic prepn;
immunomodulator epoxide peptidomimetic prepn
Peptidomimetics
   (epoxide; prepn. of epoxide peptidomimetics as irreversible HIV
   protease inhibitors)
                              40 40 de 14
Epoxides
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); USES (Uses)
   (peptidomimetics; prepn. of epoxide peptidomimetics as irreversible HIV
   protease inhibitors)
Anti-AIDS drugs
Antiviral agents
Human immunodeficiency virus 1
                 iency virus 1
Immunomodulators
   (prepn. of epoxide peptidomimetics as irreversible HIV protease inhibitors)
562-29-9P 174562+30-2P 174562-31-3P 174562-32-4P 174562-33-5P
   inhibitors)
174562-29-9P
174562-34-6P 174562-35-7P 174562-36-8P 174562-37-9P
                                                              174562-38-0P
                              174562-41-5P 174562-42-6P
                                                             174562-43-7P
174562-48-2P
174562-39-1P 174562+40-4P
                              174562-46-0P 1 174562-47-1P
174562-44-8P 174562-45-9P
                              174562-51-7P 174562-52-8P
174562-49-3P 174562-50-6P
                                                              174562-53-9P
174562-54-0P 174562-55-1P 174562-56-2P 174562-57-3P
                                                              174562-58-4P
174562-59-5P 174562-60-8P 174562-61-9P 174562-62-0P 174562-65-3P 174562-66-4P 174562-67-5P 174562-68-6P
                                                              174562-63-1P
                                                              174562-69-7P
174562-70-0P
               174562-71-1P
                               174562-72-2P
                                               174562-73-3P
                                                              174562-74-4P
174562-75-5P
               174562-76-6P
                               174562-77-7P
                                               174562-78-8P
                                                              174562-79-9P
               174562-81-3P 200262-27-7P
174562-80-2P
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); USES (Uses)
                               竹屋期 柳 明显 [4]
   (prepn. of epoxide peptidomimetics as irreversible HIV protease
                              医精乳病性三角 法人
   inhibitors)
144114-21-6, Retropepsin
RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
   (prepn. of epoxide peptidomimetics as irreversible HIV protease inhibitors)
   inhibitors)
59-67-6, 3-Pyridinecarboxylic acid, reactions 78-77-3, Isobutyl bromide
78-82-0, Isobutyronitrile 88+14-2, 2+Fürancarboxylic acid 2-Quinolinecarboxylic acid 96-41-3, Cyclopentanol 98-00-
                                                      98-00-0,
2-Furanylmethanol
                    98-59-9, paroluenesulfonyl chloride 98-98-6,
                             100-46-9, Benzylamine, reactions
2-Pyridinecarboxylic acid
                                                                100-55-0,
                   110-68-9, N-Methyl-N-butylamine 503-74-2, Isovaleric
3-Pyridylcarbinol
       527-72-0, 2-Thiophenecarboxylic acid 574-98-1,
                                                              586-98-1,
N-(2-Bromoethyl)phthalimide 586-95-8, 4-Pyridylcarbinol
                    603-35-0, Triphenylphosphine, reactions
                                                                617-89-0,
2-Pyridylcarbinol
2-Furanylmethylamine 625-45-6, Methoxyacetic acid
                                                        1113-41-3,
                  2516-33-8, Cyclopropylmethanol 4083-57-2,
L-Penicillamine
                               5163-20-2, N-Methyl-N-cyclopropylamine
3-Amino-2, 4-dimethylpentane
6921-34-2, Benzylmagnesium chloride 6964-21-2, 3-Thiopheneacetic acid
7693-46-1, p-Nitrophenyl chloroformate 13734-34-4 23844-66-8
24939-24-0, p-Aminobenzenesulfonyl chloride
                                               33445-07-7, Isopropoxyacetic
```

网络新斯特尔 网络白色 计负息通信

ST

IT

ΙT

IT

IT

ΙT

IT

59830-60-3, N#Benzyloxyčarbönyl-L-phenylalaninal 80866-93-9 96928-87-9 111491-96-4 123617-80-1, 3-Furanacetic acid 96521-86-7 136465-98-0 可加斯斯巴斯克司工 RL: RCT (Reactant) (prepn. of epoxide peptidomimetics as irreversible HIV protease inhibitors) IT 65273-64-5P 82894-53-9P 97589-56-5P 112898-22-3P 156641-79-1P 156641-83-7P 156641-81-5P 160742-44-9P 160742-45-0P 160742-70-1P 160742-71-2P 174562 82-4P 174562-83-5P 174562-84-6P 174562-85-7P 174562-89-1P 174562-90-4P 174562-88-0P 174562-86-8P 174562 92-6P 174562-94-8P 196515-98-7P 174562-91-5P 200262-29-9P 200262-28-8P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of epoxide peptidomimetics as irreversible HIV protease IT 156715-06-9P RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of epoxide peptidomimetics as lirreversible HIV protease inhibitors)

174562-91-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) ΙT (prepn. of epoxide peptidom metics as irreversible HIV protease inhibitors) 174562-91-5 HCAPLUS RN L-Valine, 3-(methylthio)-N-[(4-nitrophenoxy)carbonyl]-, phenylmethyl ester CN (9CI) (CA INDEX NAME):

Absolute stereochemistry.

ANSWER 7 OF 24

11: 1

14.00

AN1997:440050 HCAPLUS 127:66223 DN Preparation of urea molety-containing peptide derivatives as neutral ΤI endopeptidase and angiotensin converting enzyme inhibitors Nagano, Masanobu; Takenaka, Yasuhei; Kato, Takeshi IN Fujisawa Pharmaceutical Co., Ltd., Japan PAJpn. Kokai Tokkyo Koho, 51 pp. SO CODEN: JKXXAF DT Patent LA Japanese ICM C07C275-16 IC A61K031-195; A61K031-215; A61K031-445; A61K038-00; C07C275-18; C07D207-06; C07D207-16; C07D211-62; A61K031-40; C12N009-99; C07M007-00 CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 1 FAN.CNT 1 APPLICATION NO. KIND DATE PATENT NO.

HCAPLUS COPYRIGHT, 2001 ACS

L27

the fillings off.

114 1 12 12 12

1 15

```
123-75-1, Pyrrolidine, reactions 617-52-7, Dimethyl itaconate
                           Triisopropyl borate 6165-69-1, 3-Thienylboronic
     2812-46-6
                 5419-55-6
                        9000-92-4, Amylase 24424-99-5, Di-tert-butyl

0-65-8 30418-59-8, 3-Aminophenylboronic acid

gene 63024-30-6, 71989-31-6, 118786-32-6
            7693-46-1
                   28310-65-8
     dicarbonate
     32315-10-9, Triphosgene
     191426-93-4
                   191426-94-5
     RL: RCT (Reactant)
        (prepn. of urea moiety-contg. peptide derivs. as neutral endopeptidase
        and angiotensin converting enzyme inhibitors)
                  31575-75-4P 53087-13-1P 78887-39-5P
                                                             118602-51-0P
     6793-92-6P
TΤ
                                   137255-91-5P
                                                   146631-00-7P
                                                                  150351-64-7P
     118602-52-1P
                    137255-87-9P
                    190661-29-1P
                                   191425-25-9P 191425-26-0P
     156682-54-1P
     191425-27-1P
                    191425-28-2P
                                   191425-29-3P
                                                   191425-30-6P
                                                                  191425-31-7P
                                   191425-34-0P: 191425-35-1P
                                                                  191425-36-2P
     191425-32-8P
                    191425-33-9P
                                                   191425-40-8P
     191425-37-3P
                    191425-38-4P
                                   191425-39-5P
                                   191425-43-1P 191425-44-2P
                                                                  191425-45-3P
     191425-41-9P
                    191425-42-0P
                                   191425-48-6P
                                                   191425-49-7P
                                                                  191425-50-0P
     191425-46-4P
                    191425-47-5P
                    191425-52-2P
                                   191425-53-3P
                                                   191425-54-4P
     191425-51-1P
     191425-55-5P 191425-56-6P 191425-57-7P
     191425-58-8P 191425-59-9P 191425-60-2P
                                   191425-63-5P
     191425-61-3P
                    191425-62-4P
                                                   191425-64-6P
                                                                  191425-65-7P
                                  191425-68-0P 191425-69-1P 191425-70-4P
                    191425<del>,</del>67-9P
     191425-66-8P
                    191425-72-6P
                                   191425-73-7P
                                                   191425-74-8P
                                                                 191425-75-9P
     191425-71-5P
                                   191425-81-7P 191425-83-9P
                    191425-79-3P
                                                                  191425-85-1P
     191425-77-1P
                                   191425-90-8P 191425-92-0P
                    191425-88-4P
                                                                  191425-94-2P
     191425-87-3P
                                   191426-01-4P 191426-03-6P 191426-05-8P
                    191425#97-5P
     191425-95-3P
                    191426-09-2P
                                   191426-11-6P
                                                   191426-13-8P
                                                                  191426-15-0P
     191426-07-0P
                                                   191426-21-8P
                    191426 17-2P
                                   191426-18-3P
                                                                  191426-23-0P
     191426-16-1P
     191426-26-3P 191426-27-4P 191426-29-6P
                                                   191426-37-6P
     191426-31-0P
                    191426-33-2P
                                   191426-35-4P
                                   191426-41-2P 191426-42-3P
                                                                  191426-43-4P
                    191426-40-1P
     191426-39-8P
                                   191426-46-7P
                                                   191426-47-8P
                                                                  191426-48-9P
     191426-44-5P
                    191426-45-6P
                                   191426-51-4P 191426-52-5P
                                                                  191426-53-6P
     191426-49-0P
                    191426-50-3P
                                   191426-56-9P 191426-57-0P
                    191426-55-8P
                                                                  191426-58-1P
     191426-54-7P
                                    191426-61-6P
                                                   191426-62-7P
                                                                  191426-63-8P
                    191426-60-5P
     191426-59-2P
                    191426-67-2P
                                   191426-68-3P
                                                   191426-69-4P
                                                                  191426-70-7P
     191426-64-9P
                    191426-72-9P
                                   191426-73-0P
                                                   191426-74-1P
     191426-71-8P
     191426-75-2P 191426-76-3P 191426-77-4P
                                   191426-81-0P
                                                   191426-82-1P
     191426-78-5P
                    191426-80-9P
                                   1191426-85-4P
     191426-83-2P
                    191426<del>-</del>84-3P
                                                   191426-86-5P
                                   191426-89-8P
                                                   191426-90-1P
     191426-87-6P
                    191426-88-7P
                                    191426-92-3P
     191426-91-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of urea moiety-conto peptide derivs as neutral endopeptidase
        and angiotensin converting enzyme inhibitors)
IΤ
     191426-93-4
     RL: RCT (Reactant)
        (prepn. of urea molety-contg. peptide derivs. as neutral endopeptidase
        and angiotensin converting enzyme inhibitors)
                                   7种国际等的 下方下
     191426-93-4 HCAPLUS
RN
     L-Tyrosine, O-(1,1-dimethylethyl) +N+[(4-nitrophenoxy)carbonyl]-,
CN
                                    (CA INDEX NAME)
     1,1-dimethylethyl ester (9CI)
```

1

[中国 ] 118. .

The first second

Absolute stereochemistry.

1644

1 1/11

11.

11 11 11 11 11 11 11 11 11 11 11 11 11

1211年前11日 日 新特拉特 社

**新聞音舞時刊** ·清朝一年 用事止 1411年11日

1914的。1911年,

IT 191425-26-0P 191425-37-3P 191425-55-5P 191425-56-6P 191425-57-7P 191425-58-8P 191425-59-9P 191425-60-2P 191426-26-3P 191426-27-4P 191426-29-6P 191426-731-0P 191426-75-2P 191426-76-3P 191426-77-4P 191426-78-5P 191426-87-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of urea molety-contg. peptide derivs: as neutral endopeptidase and angiotensin converting enzyme inhibitors)

RN 191425-26-0 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, alpha.-[[(4-nitrophenoxy)carbonyl]amino]-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry

$$O_2N$$

H
N
S
OBu-t
Ph

RN 191425-56-6 HCAPLUS

CN 2-Benzothiazolepropanoic acid, alpha.-[[(4-nitrophenoxy)carbonyl]amino]-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191425-57-7 HCAPLUS

CN L-Tyrosine, O-(1-methylethyl)-N-[(4-nitrophenoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191425-58-8 HCAPLUS :

CN 2-Naphthalenepropanoic acid, .alpha.-[[(4-nitrophenoxy)carbonyl]amino]-, methyl ester, (S)- (9CI) (CA INDEX NAME)

RN 191425-59-9 HCAPLUS

CN 1,3-Benzodioxole-5-propanoic acid, .alpha.-[[(4nitrophenoxy)carbonyl]amino]-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191425-60-2 HCAPLUS

CN L-Tyrosine, N-[(4-nitrophenoxy)carbonyl]-O-phenyl-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191426-26-3 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, 4'-fluoro-.alpha.-[[(4-nitrophenoxy)carbonyl]amino]-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191426-27-4 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, alpha.-[[(4-nitrophenoxy)carbonyl]amino]-4'-(trifluoromethyl)-, methyl ester, (S)-(9CI) (CA INDEX NAME)

RN 191426-29-6 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, 4'-methyl-.alpha.-[[(4-nitrophenoxy)carbonyl]amino]-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

'RN 191426-31-0 HCAPLUS

CN L-Serine, O-[(4-fluorophenyl)methyl]-N-[(4-nitrophenoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191426-75-2 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, alpha.-[[(4-nitrophenoxy)carbonyl]amino]-2'-(phenylmethoxy)-, methyl ester, (S)- (9CI) (CA INDEX NAME)

RN 191426-76-3 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, .alpha.-[[(4-nitrophenoxy)carbonyl]amino]-3'-(phenylmethoxy)-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191426-77-4 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, .alpha.-[[(4-nitrophenoxy)carbonyl]amino]-4'-(phenylmethoxy)-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Horning the second

Absolute stereochemistry.

RN 191426-78-5 HCAPLUS

CN [1,1'-Biphenyl]-4-propanoic acid, 3'-(acetylamino)-alpha.-[[(4-nitrophenoxy)carbonyl]amino]-, methyl ester, (S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 191426-87-6 HCAPLUS

CN L-Lysine, N2-[(4-nitrophenoxy)carbonyl]-N6-[(phenylmethoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

ed, 50 m

```
L27 ANSWER 8 OF 24 HCAPLUS COPYRIGHT 2001 ACS
```

AN 1996:637443 HCAPLUS

DN 125:329473

TI Preparation of aminediol-containing peptide analogs as retroviral protease inhibitors

IN Gordon, Eric M.; Barrish, Joel C.; Bisacchi, Gregory S.; Sun, Chong-qing; Tino, Joseph A.; Vite, Gregory D.; Zahler, Robert

PA E. R. Squibb & Sons, Inc., USA

SO U.S., 219 pp. Cont.-in-part of U.S. Ser. No. 927,027, abandoned. CODEN: USXXAM

DT Patent

LA English

IC ICM C07D401-12

NCL 552303000

CC 34-3 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 1

FAN	•	CNT	2

2221	PATENT NO.		• •	•			
ΡI	US 5559256	A	19960924	us 1993-79978	19930625		
	AU 9341659	A1	19940127	- <sup>©</sup> HAU⊞1993-41659	19930630		
	AU 677194	B2 '	19970417	* .	:		
	HU 67090	<b>A</b> 2	19950130	HU 1993-2080	19930719		
	CA 2100894	AA	19940121:14	CA 1993-2100894	19930720		
	NO 9302620	A '	19940121	NO 1993-2620	19930720		
	EP 580402	A2	19940126	Fi EP:1993-305691	19930720		
	EP 580402	A3 :	19970305	Charles of the Secretary			
				, GB, GR, IE, IT, LI		PT,	SE
	ZA 9305243	A	19940217	1 ZA 1993-5243	19930720		
	CN 1085546	Α	19940420	CN 1993-108954	19930720		
	JP 06206857	A2	19940726	JP 1993-201016	19930720		
	US 5760036	Α.	19980602	US 1995-455295 US 1995-456125	19950531		
	US 5776933	Α	19980707	US 1995-456125	19950531		
PRAI	US 1992-916916		19920720	•			
	US 1992-927027	1.1	19920806	9 t. )			
	US 1993-79978	' i	19930625				
os	MARPAT 125:3294	73					
GI			!	,,			

Aa-E-NR8CHR9H(OH)CH2NHCH2CH(OH)CHR9NR8-E-Ab [Aa, Ab = H, alkyl, R3C(:Z),AΒ R3SO2, R3R4NSO2, R3R4NC(:Z), R3SC(:O), R5R6R7COC(:Z); E = a single bond ora peptide chain contg. 1 to 4 amino acids, the N-terminus of which is bonded to Aa or Ab; R3, R4 = H, alkyl, aryl, carbocyclyl; R5, R6, R7 = H, alkyl, aryl, carbocyclyl, fluorenyl, alkynyl, alkenyl; R5, R6, and R7 may, independently, be joined together with the carbon atom to which they are bonded, to form a mono-, bi- or tricyclic carbocyclic ring system; R8 = H, alkyl; R9 = arylalkyl; Z = O, S; wherein: wherever they appear alone or as part of another group, unless otherwise indicated, the terms "alk." or "alkyl" denote a straight or branched chain satd. radical contg. 1 to 12 carbons in the normal chain, optionally substituted by one or more groups selected from (un)protected OH, oxo (with the proviso that the carbon bearing the oxo group is not adjacent to a heteroatom), CO2H, halo, alkoxy, aryloxy, alkoxycarbonyl, etc.] or salts thereof, which inhibit retroviral protease and are particularly useful in the treatment and/or prevention of HIV infection (AIDS), are prepd. Thus, bis(3-amino-2hydroxy-4-phenylbutyl) amine deriv. (I; R = H) was condensed with L-tert-leucine deriv. (HO-Q) using 1-ethyl-3-(3dimethylaminopropyl)carbodiimide hydrochloride and HOBT in DMF/CH2CH2 at 0.degree. to room temp. to give the title compd. I (R = Q). The latter compd. at 10 .mu.M in vitro inhibited 99% HIV protease and showed IC50 of 0.012 .mu.M which was the concn. of drug that increased the formazan prodn. in CEM-SS cells infected with the RF strain of HIV to 50% of that produced by uninfected cells in the absence of drug. ST

ST aminedial contg peptide analog prepn; retroviral protease inhibitor; HIV infection AIDS treatment

IT Acquired immune deficiency syndrome Virucides and Virustats

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of aminediol-contg. peptide analogs as retroviral protease inhibitors for treatment of HIV infection (AIDS))

Peptides, preparation
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(analogs, prepn. of aminediol-contg: peptide analogs as retroviral protease inhibitors for treatment of HIV infection (AIDS))

IT Virus, animal

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (human immunodeficiency, prepn. of aminediol-contg. peptide analogs as retroviral protease inhibitors for treatment of HIV infection (AIDS)) 272-01-5P, Furo[2,3-b]pyridine 609-71-2P 2508-01-2P 3356-88-5P

3694-86-8P 7423-92-9P 13031-76-0P 15833-82-6P 15833-84-8P 22455-69-2P 27038-48-8P, Furo[2,3-b]pyridin-3(2H)-one 41036-01-5P

图1 指注扩

IT

```
52532-02-2P
               62030-47-1P
                              83096-36-0P
                                            97024-23-2P
                                                           100841-12-1P
                                                        116005-23-3P
109274-92-2P,
              Furo[2,3-b]pyridine-2-carboxaldehyde
                                               161302-40-5P
                                161302-39-2P
                                                                162537-26-0P
144731-95-3P
                161302-38-1P
                                                                162537-78-2P
                                162537-76-0P
                                                162537-77-1P
162537-37-3P
                162537-75-9P
                               162537-81-7P 162537-82-8P,
162537-79-3P
                162537-80-6P
                                  162537-83-9P
                                                  -162537-84-0P
Furo[2,3-b]pyridine-2-methanol
                               162537-88-4P
                                               162537-89-5P
                                                               162537-91-9P
                162537-87-3P
162537-86-2P
                                                               162537-99-7P
                               162537-97-5P
                                                162537-98-6P
162537-95-3P
                162537-96-4P
                                                               162538-10-5P
162538-00-3P
                162538-01-4P
                                162538-02-5P
                                               162538-08-1P
                               162538-13-8P
                                                162538-14-9P
                                                               162538-15-0P
162538-11-6P
                162538-12-7P
                162538-19-4P
                                162538-20-7P
                                                162538-21-8P
                                                               162538-22-9P
162538-18-3P
                162538-24-1P
                                162538-25-2P
                                                162538-28-5P
                                                                162538-29-6P
162538-23-0P
                162538-33-2P
                                162538-37-6P
                                                162538-38-7P
                                                                162538-39-8P
162538-31-0P
                               162538-45-6P
                                                                162538-48-9P
162538-40-1P
                162538-42-3P
                                                162538-47-8P
                               162538-52-5P
                162538-51-4P
                                               162538-53-6P
                                                                162538-54-7P
162538-50-3P
                162538-57-0P
                                162538-58-1P
                                                162538-59-2P
                                                                162538-60-5P
162538-55-8P
                162538-63-8P
                               162538-64-9P
                                               162538-65-0P
                                                               162538-66-1P
162538-61-6P
                               162538-70-7P
                                                162538-71-8P
                                                               162538-72-9P
162538-67-2P
                162538-69-4P
                               162538-76-3P
                162538-74-1P
                                                162538-80-9P
                                                                162538-81-0P
162538-73-0P
                162538-84-3P
                               162538-85-4P
                                                162538-90-1P
                                                               162538-91-2P
162538-83-2P
162538-92-3P
                162538-93-4P
                                162538-94-5P
                                                162538-95-6P
                                                                162538-96-7P
                               162539-03-9P
                                                                162539-07-3P
162538-97-8P
                162539-02-8P
                                                162539-05-1P
                               162539-15-3P
162539-09-5P
                162539-13-1P
                                               162539-17-5P
                                                                162539-18-6P
                                162539-21-1P
                                                162539-22-2P
                                                                162539-23-3P
162539-19-7P
                162539-20-0P
162539-25-5P
                162539-27-7P
                                162539-29-9P
                                                162539-32-4P
                                                                162539-33-5P
                                162539-36-8P
                                                                162539-38-0P
162539-34-6P
                162539-35-7P
                                                162539-37-9P
                                162539+43-7P 162539+44-8P
                162539-41-5P
                                                                162539-45-9P
162539-39-1P
                                                162539-50-6P
                                                                162539-54-0P
                162539-47-1P
                                162539-48-2P
162539-46-0P
                                             162539-61-9P
                                                                162539-62-0P
                162539-58-4P
                                162539-60-8P
162539-57-3P
                                162539+65-3P , 162539-66-4P
                                                                162539-67-5P
                162539#64-2P
162539-63-1P
                                                162539-71-1P
                                                                162539-72-2P
                                162539÷70-0P
162539-68-6P
                162539-69-7P
                                                162539-76-6P
                                                                162539-80-2P
162539-73-3P
                162539-74-4P
                               162539-75-5P
                162539-85-7P
                               162539-86-8P
                                                162539-87-9P
                                                                162539-88-0P
162539-81-3P
                               ·162539-91-5P
162539-89-1P
                162539-90-4P
                                                162539-92-6P
                                                                162539-93-7P
                162539-95-9P
                                162539-96-0P
                                                162539-97-1P
                                                                162539-98-2P
162539-94-8P
                162540-00-3P
                                162540-01-4P
                                                162540-03-6P
                                                                162540-04-7P
162539-99-3P
                162540-06-9P
                                162540-07-0P
                                                162540-08-1P
                                                                162540-09-2P
162540-05-8P
                               162540-12-7P
                                                162540-13-8P
                                                                162540-14-9P
162540-10-5P
                162540-11-6P
                162540-17-2P
                                162540-18-3P
                                                162540-19-4P
                                                                162540-20-7P
162540-15-0P
                162540-22-9P
                                                                162540-25-2P
                                162540-23-0P
                                                162540-24-1P
162540-21-8P
                                162540-28-5P
                                                162540-29-6P
                                                                162540-30-9P
162540-26-3P
                162540-27-4P
                                162540-33-2P
                162540-32-1P
                                                162540-34-3P
                                                                162540-35-4P
162540-31-0P
                                162540-38-7P
                                                162540-39-8P
                                                                162540-40-1P
                162540-37-6P
162540-36-5P
                                                162540-44-5P
                                                                162540-45-6P
                162540-42-3P
                                162540-43-4P
162540-41-2P
                162540-47-8P
                                162540-48-9P
                                                162540-49-0P
                                                                162540-50-3P
162540-46-7P
                                162540-53-6P
                                                162540-54-7P
                                                                162540-55-8P
162540-51-4P
                162540-52-5P
RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
   (prepn. of aminedial-contg. pertide analogs as retroviral protease
   inhibitors for treatment of HIV infection (AIDS))
                                162540-58-1P
                                                162540-59-2P
                                                                162540-60-5P
                162540-57-0P
162540-56-9P
                                                162540+66-1P
                                                                162540-67-2P
                                162540-65-0P
                162540-63-8P
162540-61-6P
                                                                162540-72-9P
                                                162540-71-8P
                               162540-70-7P
162540-68-3P
                162540-69-4P
                                                                162540-77-4P
                                                162540-76-3P
                162540-74-1P
                                162540-75-2P
162540-73-0P
                               162540-80-9P
                                                                162540-82-1P
                162540-79-6P
                                                162540-81-0P
162540-78-5P
                                162540-85-4P
                                                162540-86-5P
                                                                162540-87-6P
162540-83-2P
                162540-84-3P
162540-88-7P
                162540-89-8P 162540-90-1P
                                                162540-91-2P
                                                                162540-92-3P
                               162540-95-6P
                                               162540-96-7P
                                                                162540-97-8P
162540-93-4P
                162540-94-5P
                                                                162541-02-8P ~
                                                162541-01-7P
                162540-99-0P
                               162541-00-6P
162540-98-9P
```

直接推广 3-1957 1 m 198201 114 1-1-1-1 11-1-6-1

\*\* : : : : t

· 特别,特别: 阿拉克斯· 中华

1119時制工

11

10

TT

```
162541-03-9P
                 162541-04-0P
                                 162541-05-1P
                                                  162541-06-2P
                                                                   162541-07-3P
162541-08-4P
                 162541+09-5P
                                  162541-11-9P
                                                   162541-12-0P
                                                                   162541-13-1P
                                                                   162541-18-6P
                                  162541-16-4P
                                                   162541-17-5P
162541-14-2P
                 162541-15-3P
                                                   162541-22-2P
                                  162541-21-1P
                                                                   162541-23-3P
162541-19-7P
                 162541-20-0P
                                 162541-97-1P 162541-98-2P
                 162541-25-5P
                                                                   162541-99-3P
162541-24-4P
                                  162542-02-1P
                                                  162542-03-2P
                                                                   162542-05-4P
162542-00-9P
                 162542-01-0P
162542-08-7P
                 162542-09-8P
                                  162542-10-1P
                                                   162677-24-9P
                                                                   162677-29-4P
                                 162677-33-0P
                                                   162677-34-1P
162677-30-7P
                 162677-32-9P
                                                                   162677-35-2P
162677-36-3P
                 162677-37-4P
                                 162677-38-5P
                                                  162677-39-6P
                                                                   162677-40-9P
162677-42-1P
                 162677-45-4P
                                  162677-48-7P
                                                   162677-50-1P
                                                                   162677-52-3P
                                 162677-58-9P
                                                  162677-59-0P
                                                                   162677-60-3P
162677-54-5P
                 162677-56-7P
162677-61-4P
                 162677-62-5P
                                                   162677-64-7P
                                 162677-63-6P
                                                                   162677-65-8P
                 162677-69-2P
162677-66-9P
                                  162677-70-5P
                                                   162677-71-6P
                                                                   162677-72-7P
                                 162677-75-0P
                 162677+74-9P
                                                  162677-76-1P
                                                                   162677-77-2P
162677-73-8P
                 162677-79-4P
162677-78-3P
                                  162677-80-7P
                                                   162677-81-8P
                                                                   162677-82-9P
                                  162677-85-2P
                                                   162677-86-3P
                 162677-84-1P
                                                                   162677-87-4P
162677-83-0P
                                  162677-90-9P
                                                   162677-92-1P
                                                                   162677-93-2P
162677-88-5P
                 162677-89-6P
                                                   162677-97-6P
162677-94-3P
                 162677-95-4P
                                 162677-96-5P
                                                                   162677-98-7P
                                  162678-01-5P
                                                162678-02-6P
162677-99-8P
                 162678-00-4P
                                                                   162678-03-7P
                 162678-05-9P
                                  162678-06-0P
                                                   162678-07-1P
                                                                    162678-08-2P
162678-04-8P
                                  162678-12-8P
                                                  162678-13-9P
                                                                    162678-14-0P
162678-09-3P
                 162678-10-6P
162678-15-1P
                 162678-16-2P
                                  162678-17-3P
                                                   162678-18-4P
                                                                    162678-23-1P
162678-25-3P
                 162678-31-1P
                                  162678-33-3P
                                                   162678-34-4P
                                                                    162678-38-8P
                                                   175233-62-2P
170996-47-1P
                 170996-48-2P
                                  171228-69-6P
                                                                    175417-50-2P
                 183161-02-6P
                                  183161-31-1P
                                                   183161-35-5P
                                                                   183162-38-1P
175417-51-3P
                 183162-40-5P
183162-39-2P
                                  183162-41-6P
                                                  183162-42-7P
                                                                   183162-44-9P
                 183162-49-4P
                                  183162-51-8P
                                                   183162-58-5P
183162-45-0P
                                                                   183162-62-1P
                                                   183162-69-8P
                 183162-64-3P
                                                                   183162-75-6P
183162-63-2P
                                  183162-67-6P
                                  183255-85-8P
                                                   183255-86-9P
                 183162-79-0P
                                                                   183255-88-1P
183162-77-8P
                                  183255-92-7P
                                                                   183255-94-9P
                 183255-90-5P
                                                   183255-93-8P
183255-89-2P
183256-02-2P
RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); USES (Uses)
    (prepn. of aminediol-contg. peptide analogs as retroviral protease
   inhibitors for treatment of HIV infection (AIDS));
144114-21-6, Retropepsin
RL: BPR (Biological process); BSÜ (Biological study, unclassified); MSC
(Miscellaneous); BIOL (Biological study); PROC (Process)
    (prepn. of aminediol-contg. peptide analogs as retroviral protease
inhibitors for treatment of HIV infection (AIDS).)

50-00-0, Formaldehyde, reactions 52-52-8, 1-Aminocyclopentanecarboxylic
        56-12-2, 4-Aminobutyric acid, reactions 64-18-6, Formic acid, ns 68-12-2, Dimethylformamide, reactions 70-25-7 72-18-4,
L-Valine, reactions 74-88-4, Methyl iodide, reactions 74-89-5, Methylamine, reactions 75-16-1, Methylamagnesium bromide 75-36-5, Acetyl chloride 75-44-5, Phosgene 75-66-1, tert-Butyl mercaptan 75-98-9, Trimethylacetic acid 76-83-5, Triphenylmethyl chloride 77-76-9, 2,2-Dimethoxypropane 79-14-1, reactions 79-22-1, Methyl
                  79-44-7, Dimethylcarbamyl chloride 79-50-5,
chloroformate
                  83 33-0, 1-Indanone 91-62-3, 6-Methylquinoline
DL-Pantolactone
                           95-54-5, o-Phenylenediamine, reactions
93-10-7, Quinaldic acid
95-55-6, o-Aminophenol 96-49-1; Ethylene carbonate
                                                              98-59-9, Tosyl
            98-80-6, Phenylboronic acid 98-88-4, Benzoyl chloride
                              100-44-7; Benzyl chloride, reactions
100-39-0, Benzyl bromide
100-46-9, Benzylamine, reactions 100-52-7, Benzaldehyde, reactions
100-86-7, .alpha.,.alpha.-Dimethylphenethyl alcohol
                                                             103-74-2,
2-(2-Hydroxyethyl)pyridine
                                105-36-2, Ethyl bromoacetate
Isopropyl chloroformate 108-24-7. Acetic anhydride 109-00-2,
                      109-86-4, 2 Methoxyethanol 110-06-5, tert-Butyl
3-Hydroxypyridine
                                  · 阿爾斯 明文: hor on 1
```

AND THE RESERVE

IT

IT

110-15-6, Butanedioic acid, reactions 110-91-8, Morpholine, disulfide 111-42-2, reactions 1119-67-5, 2-Carboxybenzaldehyde reactions 122-59-8, Phenoxyacetic acid 122-98-5, 2-Anilinoethanol 2-Phenoxyethanol 137-07-5, 2-Aminothiophenol 288-32-4, Imidazole, 335-08-0, 1,1,1-Trifluoroacetone cyanohydrin 353-80-0 reactions 358-23-6, Triflic anhydride 453-20-3, 3-Hydroxytetrahydrofuran 473-85-8, 1,4-Anhydro-D-threitol 500-22-1, 3-Pyridinecarboxaldehyde 501-53-1, Benzyl chloroformate 503-38-8, Trichloromethyl chloroformate 534-03-2, 2-Amino-1, 3-propaned ol 539-74-2, Ethyl 3-bromopropionate 540-51-2, 2-Bromoethanol 541-47-9, 3,3-Dimethylacrylic acid 558-30-5, 586-98-1, 2-Pyridylcarbinol | 591-80-0, 4-Pentenoic Isobutylene oxide 593-56-6, Methoxyamine hydrochloride 594-56-9, 598-21-0, Bromoacetyl bromide 611-71-2 2,3,3-Trimethylbutene 617-94-7; Dimethylphenyl carbinol 617-35-6, Ethyl pyruvate 622-40-2, 4-(2-Hydroxyethyl) morpholine 623-08-5, 624-83-9, Methyl isocyanate 625-38-7, 3-Butenoic 2-Benzyloxyethanol N-Methyl-p-toluidine 628-41-1, 1,4-Cyclohexadiene 630-19-3, Pivalaldehyde 627-18-9 644-36-0, o-Tolylacetic acid 670-95-1, 4-Phenylimidazole 677-22-5, tert-Butylmagnesium chloride 687-47-8, te 693-89-0, 1-Methylcyclopentene 759-24-0, Diethyl L-Homo-serine (S)-Ethyl lactate tert-butylmalonate 775-06-4, DL-Meta-tyrosine 821-09-0, 4-Penten-1-ol 917-54-4, Methyllithium: 937-14-4, m-Chloroperbenzoic acid 1003-04-9 1070-83-3, tert-Butylacetic acid 1120-87-2, 4-Bromopyridine 1122-62-2-Acetylpyridine 1142-20-7 1145-80-8 1148-11-4 1149-26-4 1122-62-9, 1161-13-3 1193-47-1, 2,2-Dimethylcyclohexanone 1462-03-9, 1-Methyl-1-cyclopentanol 1609-86-5, tert-Butyl isocyanate 1664-54-6, 3-Amino-3-phenylpropionic acid 1779 #49 3, Methyltriphenylphosphonium 1826-67-1, Vinylmagnesium bromide 2018-66-8 2130-96-3 bromide 2370-61-8; 2976-75-2; (1 Naphthoxy) acetic acid 2987-16-8, 2212-75-1 3,3-Dimethylbutyraldehyde 3160-59-6 3173-56-6, Benzyl isocyanate 3240-94-6, 4-(2-Chloroethyl)morpholine 3262-72-4 3587-60-8, Benzyl 3731-51-9, 2= (Aminomethyl) pyridine 4436-24-2, chloromethyl ether 4541-32-6, 2,2-Dimethylcyclopentanone Benzyloxirane 4530-20-5 4857-04-9, 2-(Chloromethyl)benzimidazole 5034-06-0, Trimethylsulfoxonium chloride 5333-74-4, Ethyl tert-butylglyoxylate 5470-11-1, Hydroxylamine hydrochloride 6278#91-7,44-Benzyloxy-2-butanone 6290-49-9, Methyl methoxyacetate 46306-52-1, LaValine methyl ester 6351-10-6, 1-Indanol 6829-40-9, Diethyl aminomalonate hydrochloride 7326-19-4, D-Phenyllactic acid 7364-25-2, Indazolinone 7486-35-3, Vinyltributyltin 7536255+2 7677-24-9, Trimethylsilyl 7693-46-1, p-Nitrophenyl chloroformate 10147-11-2, 13031-04-4 13139-15-6 13139-16-7 13139-17-8, 3-Phenyl-1-propyne N-Benzyloxycarbonyloxy succinimide 13329+18-5, 5-Benzyloxy-2-pentanone 13570-08-6, 2-Benzimidazoleacetic acid 13575-16-1, Ethyl 5-Phenyloxazole-2-carboxylate 13734-34-4 13734-41-3 14347-78-5, (R)-2,2-Dimethyl-1,3-dioxolane-4-methanol 14397-64-9, 1-Ethoxycarbonyl-2-indanone 15761 39-4 16520-62-0, 4-Phenyl-1-butyne 17392 83 5, (R) -Methyl lactate 17463-43-3, 16677-29-5 17199-29-0 DL-3,3,3-Trifluoroalanine 18162-48-6, tert-Butyldimethylsilyl chloride 19575-07-6, Methyl quinaldate 19728-63-3, Z-Thr-OH 18942-49-9 19752-84-2, 3-Hydroxytetrahydropyran 19810-31-2, Benzyloxyacetyl 20117-47-9, 1-Methyl-1-cyclobutanol 20160-60-5, 2-Trimethylsilylethyl chloroformate 20412-38-8, Neopentyl chloroformate 20662-89-9, 4-Phenyloxazole 20859-02-3; L-tert-Leucine 21641-92-9 22323-82-6, (S)-2,2-Dimethyl-1,3-dioxolane-4-methanol 22146-57-2 24424-99-5, Di-tert-butyl dicarponate 26628-22-8, Sodium azide 26782-71-8, D-tert-Leucine 28920-43-6, 9-Fluorenylmethyl chloroformate 29943-42-8, Tetrahydro-4H-pyran-4-one 28954-12-3, L-Allothreonine 30525-89-4, Paraformaldehyde 32366-02-2, N-Benzyl-N-methyl carbamoyl 36024-28-9 37595-74-7, N-Phenyltriflimide 37736-82-6, chloride

```
40299-87-4,
N-tert-Butoxycarbonyl-L-cyclohexylalanine
4-(Bromoacetyl)morpholine 41242-94-8, 2-Hydroxymethyl quinoxaline
52373-72-5 53333-76-9, 2,2-Dimethy1-1-propanesulfonyl chloride 58632-95-4, Boc-ON 59562-82-2 60456-21-5 67478-50-6 6883
                                                                       68835-89-2,
                                69739-34-0 tert-Butyldimethylsilyl triflate
Di-tert-amyl dicarbonate
76513-69-4, 2-(Trimethylsilyl) ethoxymethyl chloride 78879-20-6
                               86087\frac{1}{2}23-2, (S)-\frac{1}{2}(+)-3-Hydroxytetrahydrofuran
               85613-64-5'
80360-23-2
                                 112372-06-2, Furo[2,3-c]pyridine-2-
                107202-43-7
106167-47-9
                    127862-89-9 162537-72-6, Furo[2,3-c]pyridine-2-methanol
carboxaldehyde
                                               162870-63-5
                                 162678-30-0
162537-73-7
                162541-63-1
                                            70 FT
RL: RCT (Reactant)
    (prepn. of aminediol-contg. peptide analogs as retroviral protease
   inhibitors for treatment of HIV infection (AIDS))
95-13-6P, 1H-Indene | 102-14-7P | 111-32-0P
                                                      272-62-8P,
Furo[3,2-b]pyridine 334-88-3P, Diazomethane
                                                       374-35-6P
                                          1191-31-7P 1615-14-1P,
              815-17-8P
                            1184-93-6P
587-33-7P
1H-Imidazole-1-ethanol 1780-17-2P, 2-Quinolinemethanol
                                                                      1796-25-4P
               2258-42-6P, Acetic formic anhydride 2280-28-6P
                                                                            2644-82-8P
2215-63-6P
               2849-93-6P, 1H-Benzimidazole-2-carboxylic acid
                                                                          3587-64-2P
2842-44-6P
3724-55-8P 4026-20-4P 4113-04-6P, 6-Quinolinecarboxaldehyde

4441-30-9P, 4-Morpholinepropanol 4647-42-1P 4647-43-2P 4754-2'

4856-97-7P, 1H-Benzimidazole-2-methanol 5105-78-2P 5367-24-8P

6970-72-5P 7467-35-8P 7525-64-6P 7748+36-9P, 3-Oxetanol

13737-35-4P 1447-66-8P 14598-96-0P 145546-08-4P 17450-34-9P
                                                                         4754-27-2P
18096-68-9P, 1H-Indene-2-methanol 19458+29-8P 19539-50-5P, Furo[2,3-c]pyridine 20120-24-5P 20361-09-5P 22892-29-1P
22929-52-8P 23249-97-0P, 1H-Benzimidazole-2-propanoic acid 24621-70-3P, 1H-Indole-2-methanol 25854-85-7P 25854-87-9P 30293-86-8P 31562-43-3P 33905-47-4P 34637-40-6P 35677
                                                                           24580-44-7P
                                                                  35677-88-4P
                 37859-42-0P, 2-Benzothiazolemethanol 39497-64-8P
37535-57-2P
                           2P 50411-26-2P 50531-59-4P 51110-97-5P, 53346-03-5P, 56365-70-9P 57443-39-7P
                 42417-65-2P
40594-83-0P
2-Benzoxazolepropanol!
                                 60651-97-0P: 1 62565-28-0P 62965-10-0P
                 60398-41-6P
59524-02-6P
                                 67706 \pm 63 \pm 2P, 170448 \pm 03 \pm 2P, 73282 \pm 11 \pm 8P
64360-69-6P
                 66866-64-6P
77186-95-9P, 2-Benzokazolemethanol 80466-51-9P 85328-36-5P
                                 86096-65-3P 86562-71-2P
                                                                  ,88246-12-2P
85951-09-3P
                 85995-53-5P
89464-59-5P
                                 91968-72-8P = 94882-74+3P
                                                                   98737-29-2P
                 90819-30-0P
                                 98997-01-4P + 100516+88-9P,
                 98955-64-7P
98760-08-8P
6-Quinolinemethanol | 100868-72-2P | 102123-74-0P | 102123-85-3P
                                   102831-44-7P 104948-22-3P
                  102229-10-7P
                                                                       106513-42-2P
102152-03-4P
                  112372 05-1P, Furo [3,2-b] pyridine-2-carboxaldehyde
108957-20-6P
                  113459-50-0P
                                   1114645-18-0P 115916-75-1P
                                                                       127041-02-5P
113247-51-1P
                                  131424-20-9P 134807+06-0P
                  128018-44-0P
                                                                       134807-20-8P
127382-65-4P
                                   134807-30-0P 1 137515-66-3P 138432-95-8P
                 134807-29-7P
134807-28-6P
                                   1348077372446-7P 1443372-4778P 143576-95-8P 144186-52-7P 144825-44-5P 149357-61-9P 154612-75-6P 162536-40-5P 162536-41-6P 162536-46-1P
141978-97-4P
                  143372-45-6P
                  144186-00-5P
143688-65-7P
153291-20-4P
                  154117-17-6P
                  160232-54-2P
159259-43-5P
                                   162536-44-9P 162536-45-0P
162536-42-7P
                  162536-43-8P
                                                                       162536-46-1P
                                   162536-50-7P 162536-54-1P
                  162536 48-3P
162536-47-2P
                                                                        162536-55-2P
                                   162536-58-5P 162536+59-6P
                                                                       162536-60-9P
162536-56-3P
                  162536-57-4P
                                   162536-65-4P 162536+67-6P
                 162536-64-3P
                                                                       :162536-68-7P
162536-63-2P
                                    162536-71-2P
                                                     162536-72-3P
                                                                       162536-73-4P
                  162536-70-1P
162536-69-8P
                                    162536-78-9P 162536-79-0P
                                                                        162536-80-3P
                  162536-77-8P
162536-74-5P
                                    162536-83-6P 11 162536-84-7P
                                                                        162536-85-8P
162536-81-4P
                  162536-82-5P
                                    162536-88-1P 162536-89-2P
                                                                        162536-90-5P
162536-86-9P
                  162536-87-0P
                                    162536-93-8P + 162536-94-9P
                                                                       162536-95-0P
                  162536-92-7P
162536-91-6P
                                    162536-98-3P 162536-99-4P
                                                                        162537-00-0P
162536-96-1P
                  162536-97-2P
                  162537+03-3P 162537-10-2P 162537-11-3P
162537-01-1P
                                    162537-14-6P 162537-15-7P
                                                                       '162537-16-8P
                  162537-13-5P
162537-12-4P
```

APPLEASE REPORTED

4

ΙT

```
162537-21-5P
                    162537-20-4P
                                                   162537-22-6P
                                                                  .162537-27-1P
     162537-17-9P
                                                   162537-34-0P
                    162537-32-8P
                                    162537-33-9P
                                                                   162537-35-1P
     162537-31-7P
                                                   162537-41-9P
                                                                  162537-42-0P
                    162537-39-5P
                                    162537-40-8P
     162537-36-2P
                    162537-44-2P
                                   162537-45-3P
                                                   162537-46-4P
                                                                   162537-47-5P
     162537-43-1P
                                    162537-50-0P
                                                   162537-51-1P
                                                                   162537-53-3P
     162537-48-6P
                    162537-49-7P
                                   :162537-56-6P
                                                  +162537-61-3P,
     162537-54-4P
                    162537-55-5P
     Furo[3,2-b]pyridine-2-methanol
                                     #162537-62-4P
                                                     162537-63-5P
                                    162537-66-8P 162537-67-9P
                                                                   162537-68-0P
                    162537-65-7P
     162537-64-6P
                                   162537-74-8P
     162537-69-1P
                    162537#70-4P
                                                 ii.
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of aminediol-contg. peptide analogs as retroviral protease
        inhibitors for treatment of HIV infection (AIDS))
                                                                   162538-17-2P
                    162538-07-0P
                                   162538-09-2P
                                                   .162538-16-1P
IT
     162538-06-9P
                                   162538-34-3P 162538-36-5P
                                                                   162538-41-2P
     162538-30-9P
                    162538-32-1P
                                                  162538-49-0P
                                    162538-46-7P 162538-82-1P
                    162538-44-5P
                                                                   162538-56-9P
     162538-43-4P
                    162538-77-4P
                                                   162538-86-5P
                                                                   162538-98-9P
     162538-68-3P
                    162539-00-6P
                                    162539-01-7P
                                                   162539-04-0P
                                                                   162539-06-2P
     162538-99-0P
                                   162539-12-0P
                    162539-11-9P
                                                   162539-14-2P
                                                                   162539-16-4P
     162539-08-4P
                                    162539-28-8P 162539-30-2P
                                                                   162539-31-3P
     162539-24-4P
                    162539-26-6P
                                                  162539+52+8P
                    162539-42-6P
                                   162539-51-7P
                                                                   162539-53-9P
     162539-40-4P
                    162539-56-2P
                                    162539-77-7P
                                                   162539-78-8P
                                                                   162539-79-9P
     162539-55-1P
                                                  162541-29-9P
                                   162541-28-8P
                                                                   162541-30-2P
                    162541-27-7P
     162540-62-7P
                                    162541-33-5P
                                                   162541-34-6P
                                                                   162541-35-7P
     162541-31-3P
                    162541-32-4P
                                                   162541-41-5P
                    162541-39-1P
                                    162541-40-4P
                                                                   162541-42-6P
     162541-38-0P
                                   162541-45-9P
     162541-43-7P
                    162541-44-8P
                                                   162541-46-0P
                                                                   162541-48-2P
                                                   162541-53-9P
                                    162541-52-8P
                                                                   162541-54-0P
     162541-49-3P
                    162541-50-6P
                                    162541-58-4P
                                                  162541-60-8P
                                                                   162541-61-9P
                    162541-57-3P
     162541-55-1P
                                    162541-65-3P 162541-66-4P
                                                                   162541-67-5P
                    162541-64-2P
     162541-62-0P
                                    162541-70-0P
                    162541-69-7P
                                                   162541-71-1P
                                                                   162541-72-2P
     162541-68-6P
                                    162541÷75-5P
                                                   162541-76-6P
                                                                   162541-77-7P
                    162541-74-4P
     162541-73-3P
                                                   162541-81-3P
162541-87-9P
                    162541-79-9P
                                   162541-80-2P
                                                                   162541-82-4P
     162541-78-8P
                    162541-85-7P
                                    162541-86-8P
                                                                   162541-88-0P
     162541-84-6P
                                   162541-94-8P
                                                   162541-95-9P
                                                                   162541-96-0P
                    162541-92-6P
     162541-89-1P
                                                . 162677-41-0P
                                   162677-31-8P
                                                                   162677-43-2P
                    162677-23-8P
     162677-20-5P
                                   162677-49-8P
                                                   162677-51-2P
                                                                   162677-53-4P
                    162677-47-6P
     162677-46-5P
                                    162677÷91-0P
                                                   162678-21-9P
                                                                   162678-22-0P
     162677-55-6P
                    162677<del>-</del>57-8P
                                   162678-36-6P
                                                   162678-37-7P
                                                                   162776-41-2P
                    162678-29-7P
     162678-26-4P
                    165727-45-7P
                                    170996-45-9P
                                                   171230-81-2P
                                                                   175390-83-7P
     165524-61-8P
     183161-49-1P
                    183161-50-4P
                                    183161-56-0P
                                                   183161-57-1P
                                                                   183161-58-2P
     183161-59-3P
                                    183161-61-7P
                                                   183161-62-8P
                                                                   183161-63-9P
                    183161-60-6P
                    183161-65-1P
                                   183161-67-3P
                                                   183161-68-4P
                                                                   183161-69-5P
     183161-64-0P
                    183161-72-0P
                                    183161-73-1P
                                                   183161-74-2P
                                                                   183161-75-3P
     183161-71-9P
                    183161-77-5P
                                   183161-78-6P
     183161-76-4P
                                                   183161-81-1P
                                                                   183161-82-2P
                                                   183161-86-6P
                    183161-84-4P
                                    183161-85-5P
                                                                   183161-87-7P
     183161-83-3P
                    183161-90-2P
                                    183161-91-3P
                                                   183161+92-4P
                                                                   183161-93-5P
     183161-89-9P
                    183161-95-7P
                                    183161-96-8P
                                                   183161-97-9P
                                                                   183161-98-0P
     183161-94-6P
                    183162-00-7P
                                    183162-01-8P
                                                   183162-02-9P
                                                                   183162-03-0P
     183161-99-1P
                                                                   183162-08-5P
                                    183162-06-3P
                    183162-05-2P
                                                   183162-07-4P
     183162-04-1P
                                   183162-11-0P
                                                   183162-12-1P
                                                                   183162-13-2P
     183162-09-6P
                    183162-10-9P
                                                   183162-17-6P
                                    183162-16-5P
                                                                   183162-18-7P
                    183162-15-4P
     183162-14-3P
                                   183162-21-2P
                                                   183162-22-3P
                                                                   183162-23-4P
     183162-19-8P
                    183162-20-1P
                                   183162-26-7P
                                                   183162-27-8P
                                                                   183162-28-9P
                    183162-25-6P
     183162-24-5P
                                   183162-32-5P
                                                                   183162-34-7P
                    183162-30-3P
                                                   183162-33-6P
     183162-29-0P
                    183162-36-9P
                                   183162-37-0P
                                                   183255-95-0P
                                                                   183255-96-1P
     183162-35-8P
                                   183255-99-4P
                    183255-98-3P
                                                  183256-00-0P
                                                                   183256-01-1P
     183255-97-2P
                                   183256-06-6P
                                                   183256-07-7P
                                                                   183256-08-8P
                    183256-05-5P
     183256-04-4P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of aminedial-contg. peptide analogs as retroviral protease
        inhibitors for treatment of HIV infection (AIDS))
                                     智力技能多
```

我要帮助

. 1

1111

3 3 7 1 1

ΙT

```
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of aminediol-contg. peptide analogs as retroviral protease inhibitors for treatment of HIV infection (AIDS))
162537-10-2 HCAPLUS
L-Valine, N-{(4-nitrophenoxy) carbonyl}, methyl ester (9CI) (CA INDEX NAME)
```

Absolute stereochemistry:

RN

CN

```
ANSWER 9 OF 24 HCAPLUS COPYRIGHT 2001 ACS
L27
    1996:551382 HCAPLUS
ΑN
DN
    125:196378
TΙ
    Nitrogen mustard prodrugs with hovel lipophilic protecting groups, and
    processes for their production
Springer, Caroline Joy; Niculescu-Duvaz, Ion
IN
    Cancer Research Campaign Technology Limited, UK
PA
                                hnology Limb
SO
    PCT Int. Appl., 44 pp
    CODEN: PIXXD2
DΤ
    Patent
    English
LA
    ICM C07C275-32
IC
    ICS C07C271-54; C07C219-34; C07C271-26; A61K031-17; A61K031-27
    34-2 (Amino Acids, Peptides, and
CC
    Section cross-reference(s): 1, 25, 63
FAN.CNT 1
                                        APPLICATION NO.
    PATENT NO.
                     KIND
                           DATE
                                    WO 1996-GB112
                     A1
                           19960725
                                                          19960119
ΡI
    WO 9622277
        W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE,
            ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT,
            LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE,
            SG, SI
        RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE,
            CA 2210347
    AU 9644535
    AU 709251
    EP 804413
                      B1
                          20000517
    EP 804413
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE
                     T2 19981202 JP 1996-522127 E 20000615 AT 1996-900627
                                                          19960119
    JP 10512565
                                                          19960119
    AT 193012
                    T3 20001101
    ES 2149445
                                        ES 1996-900627
                                                          19960119
                          19991221
                                         US 1997-875099
                                                          19970716
    US 6005002
                      A:
                     A V
                          19950119
PRAI GB 1995-1052
                           19960119
    WO 1996-GB112
    MARPAT 125:196378
OS
GΙ
```

**企业的 拉斯斯** 

```
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
                                              The invention provides compds. I and II [X, Y = Cl, Br, iodo, CH3SO3, or
AB
     OSO2Ph (wherein Ph is optionally substituted by 1-5 alkyl, halo, cyano,
     and/or nitro); R1, R2 each = 1-4 optional substituents; Z1, Z2 = O, NH; R3
     = H, t-Bu, allyl; Z3 = hydrocarbyl group such as carboxyethyl, optionally
     contg. heteroatoms] and their physiol. acceptable derivs. The compds. can
     be converted in situ into nitrogen mustard agents by the actions of
     enzymes such as carboxypeptidase or nitroreductase and are useful for the
     treatment of cancer (no data). For example, the glutamate ester
     L-Me3COCOCHRCH2CH2CO2CMe3 [III; R = NH2] was converted to the isocyanate
     III [R = NCO], which reacted with 4-hydroxybenzaldehyde to give 72% III [R
     = NHCO2C6H4CHO-4]. This was reduced with NaBH3CN to give 86% of III [R =
     NHCO2C6H4CH2OH-4], which was coupled with the nitrogen mustard
     4-(OCN)C6H4N(CH2CH2Cl)2 in 90% yield, followed by deesterification with
     formic acid (87%), to give title compd. IV.
ST
     glutamate nitrogen mustard prodrug antitumor prepni
ΙT
     Neoplasm inhibitors
        (prepn. of lipophilic glutamate-based nitrogen mustard prodrugs as
                                    anticancer agents):
ΙT
     Amino acids, preparation
     Amino acids, preparation
Nitrogen mustards
RL: BAC (Biological activity or effector, except adverse); RCT (Reactant);
     SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
        (prepn. of lipophilic glutamate-based nitrogen mustard prodrugs as
                                   anticancer agents)
IT
     Therapeutics
        (pharmaco-, GDEPT (gene-directed enzyme prodrug therapy); prepn. of
        lipophilic glutamate-based nitrogen mustard prodrugs as anticancer
     agents)
Pharmaceutical dosage forms
ΙT
        (prodrugs, prepn. of lipophilic glutamate-based nitrogen mustard
        prodrugs as anticancer agents)
     9031-98-5, Carboxypeptidase 9037#41-6, Nitroreductase
IT
     RL: BSU (Biological study, unclassified); MSC (Miscellaneous); BIOL (Biological study)
     (Biological study)
                                   ្នាត់ ស្រាស់ម៉ូត
        (activation by; prepn. of lipophilic glutamate-based nitrogen mustard
        prodrugs as anticancer agents):
IT
     18483-99-3P, 4-Nitrobenzyl 2-tetrahydropyranyl ether
                                                             18484-05-4P,
     4-Aminobenzyl 2-tetrahydropyranyl ether 113068-95-4P, 4-Isocyanatobenzyl
     tert-butyldiphenylsilyl ether 20161803-03-8P, 4-Nitrobenzyl
     tert-butyldiphenylsilyl ether 161803-04-9P, 4-Aminobenzyl tert-butyldiphenylsilyl ether 161803-05-0P 161803-06-1P 180839-06-9P, 4-[N,N-Bis(2-chloroethyl)amino]phenyl trimethylsilyl ether
                                  180839-12-7P 180839-13-8P 180839-14-9P 180839-17-2P 180839-18-3P 180839-19-4P
                    180839-11-6P
     180839-08-1P
     180839-15-0P
                    180839-16-1P
                                 180841-62-7P
     180839-20-7P 180839-21-8P
     RL: RCT (Reactant); SPN (Synthètic preparation); PREP (Preparation)
        (intermediate; preph. of lipophilic glutamate-based nitrogen mustard
        prodrugs as anticancer agents)
                                  180839-02-5P 180839-04-7P
                    180839-00-3P
IT
     180838-97-5P
     RL: BAC (Biological activity or effector, except adverse); RCT (Reactant);
     SPN (Synthetic preparation); THU! (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
        (prepn. of lipophilic glutamate-based nitrogen mustard prodrugs as
                                   anticancer agents):
                                   180839-03-6P 180839-05-8P
     180838-98-6P
                    180839-01-4P
ΙT
                                   in the line of the supper
```

1 1

16 41 11

RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of lipophilic glutamate based nitrogen mustard prodrugs as anticancer agents)

er agents)
123-08-0, 4-Hydroxybenzaldehyde 619-73-8, 4-Nitropenzyl
23-08-0, 4-Hydroxybenzyl
623-05-2, 4-Hydroxybenzyl
7693-46-110-87-2 ΙT 623-04-1, 4-Aminobenzyl alcohol 623-05-2, 4-Hyd 1204-69-9, 4-[N,N-Bis (2-chloroethyl) amino]phenol alcohol alcohol 20845-16-3 4-Nitrophenyl chloroformate 32677-01-3, Di-tert-butyl 57529-05-2, 4-(1,3-Dithian-2-yl)phenol L-glutamate hydrochloride 58479-61-1, tert-Butyldiphenylchlorösilane 82484-59-1, 4-[N, N-Bis (2-chloroethyl) amino] phenyl isocyanate RL: RCT (Reactant)

(starting material; prepn. of lipophilic glutamate-based nitrogen mustard prodrugs as anticancer agents)

IT 180839-20-7P 180839-21-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (intermediate; preph. of lipophilic glutamate-based nitrogen mustard prodrugs as anticancer agents) 🔠 🍀

180839-20-7 HCAPLUS RN

CN L-Glutamic acid, N-[[4-[[[(4-nitrophenoxy)carbonyl]oxy]methyl]phenoxy]carb onyl]-, bis(1,1-dimethylethyl) ester (9CI); (CA INDEX NAME)

Absolute stereochemistry.

180839-21-8 HCAPLUS RN

CN L-Glutamic acid, N-[[4-[[[(4-nitrophenoxy)carbonyl]oxy]methyl]phenoxy]carb onyl]-, di-2-propenyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

Сн2

```
COPYRIGHT 2001 ACS
    ANSWER 10 OF 24 HCAPLUS
L27
     1996:509810 HCAPLUS
AN
     125:168650
DN
    High-purity N-[4-[N,N-bis(2-iodoethyl)amino]phenoxycarbonyl]-L-glutamic
ΤI
     acid and hydrogen iodide salt as prodrugs for ADEPT therapy
IN
    Heaton, David William; Dines, Susan; Dowell, Robert Ian
PA
     Zeneca Limited, UK
so
     PCT Int. Appl., 18 pp.
     CODEN: PIXXD2
DT
    Patent
    English
LA
IC
    ICM C07C271-54
     ICS A61K031-325
CC
     34-2 (Amino Acids, Peptides, and
     Section cross-reference(s): 1, 63
FAN.CNT 1
                      KIND DATE
     PATENT NO.
                                          WO 1995-GB2997
                            19960704
                                                             19951221
ΡI
    WO 9620169
            AL, AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES,
             FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG,
             SI, SK
         RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE,
             IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR,
                                   the out of some in his
             NE, SN, TD, TG
                                        ZA 1995-10936
                            19960624
     ZA 9510936
                                                             19951221
                       Α
                                           CA 1995-2204741
                            19960704
                                                             19951221
    CA 2204741
                       AA:
                                         AU 1996-42700
    AU 9642700
                       A1
                            19960719
                                                             19951221
    AU 702337
                       B2 :
                            19990218
                                         GB 1997-8977
                            19970716
                                                             19951221
    GB 2309031
                       A1 .
                                         GB 2309031
                       B2
                            19980812
                                                             19951221
    EP 799193
                       A1
                            19971008
                                           EP 1995-941219
    EP 799193
                       B1 14 20000816
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE
    CN 1171100
                       A 19980121
                                           CN 1995-196993
                                                             19951221
                       A2 19980330
    HU 77288
                                          HU 1997-2207
                                                             19951221
                       A. 1, 19980609; . 4; 1988; 1995-10463
                                                             19951221
    BR 9510463
                            20000915 AT 1995-941219
                                                             19951221
    AT 195511
                                           FI 1997-2630
     FI 9702630
                            19970618<sup>-</sup>
                                                             19970618
                       A 1 19970620 NO 1997-2883.
    NO 9702883
                                                             19970620
                       19970623
    US 5981791
                       A 19941223
PRAI GB 1994-26133
    WO 1995-GB2997
                            19951221
    MARPAT 125:168650
os
GΙ
```

A new salt of the prodrug N-[4-[N,N-bis(2-iodoethyl)amino]phenoxycarbonyl]-AΒ L-glutamic acid (I), which is useful in antibody directed enzyme prodrug therapy (ADEPT), is disclosed. This salt, the hydrogen iodide salt I.HI, is obtained in cryst, form with m.p. 142-145.degree. and has a specified X-ray powder diffraction spectrum. Prepn. of cryst. I.HI enables I to be prepd. in a highly pure form. I.HI also has advantages of good thermal stability, easy synthesis, and reduced hygroscopic/deliquescence properties. Improved stability may be due to reversal of degrdn. of the mustard alkylating arms of the mol. by HI. A multi-step prepn. of I from L-glutamic acid and p-02NC6H4OCOCl is given. Mouse tumor xenograft expts. using therapy with F(ab')2A5B7-CPG2 conjugate (2.5 mg/kg i.v.) and I.HI (3 doses at 70 mg/kg i.p. each) gave tumor regression and >30 day growth delays with only minor effects on body wt. and peripheral white blood cell

iodoethylaminophenoxycarbonylglutamic acid hydriodide antitumor prodrug ST ADEPT; bisiodoethylaminophenoxycarbonylglutamic acid prepn antitumor prodrug ADEPT

Neoplasm inhibitors IT

(prepn. of high-purity [[bis(iodoethyl)amino]phenoxycarbonyl]glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT)

ΙT Pharmaceutical dosage forms

(prodrugs, antibody-directed enzyme prodrug therapy (ADEPT); prepn. of high-purity [[bis(iodoethyl)amino]phenoxycarbonyl]glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT) 1 ! .

Pharmaceutical dosage forms IT

(prodrugs, prepn. of high-purity [[bis(iodoethyl)amino]phenoxycarbonyl] glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT)

5269-43-2P, L-Glutamic acid bis(trimethylsilyl) ester 156079-00-4P 156079-89-9P 172974-17-3P 180031-70-3P 156079-01-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (intermediate; prepn. of high-purity [[bis(iodoethyl)amino]phenoxycarbo nyl]glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT)

电影电影电影 人名拉

IT 180031-69-0P

ΙT

IT

RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); IMF (Industrial manufacture); PRP (Properties); PUR (Purification or recovery); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of high-purity [[bis(iodoethyl)amino]phenoxycarbonyl]glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT)

156079-88-8P

RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); IMF (Industrial manufacture); PUR (Purification or recovery); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

19 1 2 1 E 2 1 S

(prepn. of high-purity [[bis(iodoethyl)amino]phenoxycarbonyl]glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT) 

4121

Artonia 4 15

IT 56-86-0, L-Glutamic acid, reactions 75-21-8, Ethylene oxide, reactions 75-77-4, Chlorotrimethylsilane, reactions 115-11-7, Isobutylene, reactions 7693-46-1, p-Nitrophenyl chloroformate RL: RCT (Reactant) (starting material; prepn. of high-purity [[bis(iodoethyl)amino]phenoxy

(starting material; prepn. of high-purity [[bis(iodoethyl)amino]phenoxy carbonyl]glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT)

IT 156079-00-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (intermediate; prepn. of high purity [[bis(iodoethyl)amino]phenoxycarbo nyl]glutamic acid and hydriodide salt as antitumor prodrugs for ADEPT)

RN 156079-00-4 HCAPLUS

CN L-Glutamic acid, N-[(4-nitrophenoxy) carbonyl]-, bis(1,1-dimethylethyl) ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L27 ANSWER 11 OF 24 HCAPLUS COPYRIGHT 2001 ACS

AN 1996:171803 HCAPLUS

DN 124:233139

Preparation of sulfonylamino acid amides containing cis-epoxide as irreversible HIV protease inhibitors in Yoon, Heungsik; Choy, Nakyen; Kim, Sung Chun; Choi, Ho II; Son, Young

IN Yoon, Heungsik; Choy, Nakyen; Kim, Sung Chun; Choi, Ho II; Son, Young Chan; Park, Chi Hyo; Moon, Kwang-Yul; Jung, Wonhee; Kim, Chung Ryeol; et al.

15 1

PA IG Chemical Ltd., S. Korea

SO Eur. Pat. Appl., 58 pp. CODEN: EPXXDW

DT Patent

LA English

IC ICM C07D303-36

ICS C07D303-40; C07D405-12; C07D407-12; A61K031-335; A61K031-435

CC 34-2 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 1

FAN.CNT 7

APPLICATION NO. KIND: DATE DATE A2 19951220 EP 1995-108908 PΙ EP 687675 19950609 687675 A3 19960306 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE 125117 B1 19971205 KR 1994-13423 19940615 EP 687675 KR 125117 A2 19960730 JP 1995-172733 JP 08193077 B2产 19991206 学 科特法 前提出 JP 2987313 A 19940615 PRAI KR 1994-13423 MARPAT 124:233139 OS GΙ

Novel cis-epoxide compds. [I; R1, R2 = H, alky1; R3 = (un) substituted arylor alky1; R4 = H, C1-4 alky1; n = 0,1,2; A = (X) (Y) mR5, NR6R7, ZCHR8R9; AΒ wherein X = CO, COCO, CO, COCO, COCOC, COCO, COCO, COCO, COCO, COCO, COCO, COCO, COCheterocyclyl, straight or branched or cyclic C1-8 alkyl or alkoxy, heterocyclylalkyl, cycloalkylalkyl, arylalkoxy; R6 = straight or branched C1-8 alkyl, cycloalkyl, cycloalkylalkyl; R7 = H, alkyl; Z = O, NH, NMe; R8, R9 = alkyl optionally substituted by arom. hydrocarbyl or cycloalkyl, C3-8 cycloalkyl, aryl], useful for treating or preventing diseases caused by HIV infection, are prepd. The novel HIV protease inhibitor I has a specific structure to form a stable bonding with the enzyme active site, which entails a highly enhanced irreversible inhibition against HIV protease. An anti-AIDS or immunomodulator contains a therapeutically effective amt. of said cis-epoxide I. Thus, (S)-5-[(Nbenzyloxycarbonyl)amino]-6-phenyl-(cis)-3-hexene-1-carboxylic acid was condensed with (S)-2-amino-3-methyl-1-phenylbutane using N-ethyl-N'-(3-(dimethylamino)propyl)carbodiimide hydrochloride (EDC) and HOBT in DMF followed by epoxidn. with m-chloroperbenzoic acid in CH2Cl2 to give the cis-epoxide, (II; R = PhCH2O2C), which was hydrogenolyzed in the presence of 10% Pd-C in MeOH under an atm. of H, coupled with  $\label{eq:n-benzyloxycarbonyl-.beta.-(S-methyl)-L-valine using EDC and HOBT in DMF \ ,$ and oxidized with m-chloroperbenzoic acid in CH2Cl2 to give the title compd. II (R = Q). The latter compd. in vitro inhibited HIV protease with the inhibition const. Kina/Ki min-1M-1 109-1010 (Kina = a rate const. indicating rate of chem. reaction forming covalent bond between an enzyme and an inhibitor in Michaelis-Menten complex; Kir = an inhibition const. indicating the dissoch. rate of Michaelis-Menten complex into an enzyme and an inhibitor) and in vitro showed IC50 of 1 nM for inhibiting the HIV-1 infection of H9 or Sup Tlacell lines. cis epoxide prepn HIV protease inhibitor; irreversible HIV protease ST

inhibitor; sulfonylamino acid amide contg cis epoxide

Acquired immune deficiency syndrome IT 94 g 3 2 34 34 34 3 Virucides and Virustats

(prepn. of sulfonylamino acid amides contg. cis-epoxide as irreversible HIV protease inhibitors for treating AIDS)

IT Amides, preparation

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(amino, prepn. of sulfonylamino acid amides contg. cis-epoxide as irreversible HIV protease inhibitors for treating AIDS)

ΙT Epoxides

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic 我们的是好,事的一个

Maria H Baltell Leinin

基础的 的复数 计图象

```
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
        (Preparation); USES (Uses)
              (cis-, prepn. of sulfonylamino acid amides contg. cis-epoxide as
             irreversible HIV protease inhibitors for treating AIDS) us, animal
IT
              (human immunodeficiency 1, prepn. of sulfonylamino acid amides contg.
             cis-epoxide as irreversible HIV protease inhibitors for treating AIDS)
                                 174562-30-2P 174562-31-3P 174562-32-4P
                                                                                                               174562-33-5P
IT
        174562-29-9P
        174562-34-6P 174562-35-7P 174562-36-8P 174562-37-9P 174562-39-1P 174562-40-4P 174562-41-5P 174562-42-6P
                                                                                                               174562-38-0P
                                                                                                             174562-43-7P
                                                           174562-41-5P 174562-42-6P 174562-47-1P
                                                                                                               174562-48-2P
        174562-44-8P 174562-45-9P
                                                           174562-51-7P 174562-52-8P 174562-53-9P 174562-56-2P 174562-57-3P 174562-58-4P 174562-63-1P 174562-63-1P
        174562-49-3P 174562-50-6P
        174562-54-0P 174562 55-1P
        174562-59-5P 174562-60-8P
                                                           174562-66-4P 174562-67-5P
                                                                                                               174562-68-6P
        174562-64-2P 174562-65-3P
                                                           174562-71-1P
                                                                                     174562-72-2P
        174562-69-7P
                                 174562-70-0P
                                                                                                               174562-73-3P
                                                          174562-76-6P
                                                                                     174562-77-7P
                                 174562-75-5P
                                                                                                               174562-78-8P
        174562-74-4P
                                                          174562-81-3P
        174562-79-9P
                                 174562-80-2P
        RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
        preparation); THU (Therapeutic use); BIOL (Biological study); PREP
        (Preparation); USES (Uses)
             (prepn. of sulfonylamino acid amides contg. cis-epoxide as irreversible
             HIV protease inhibitors for treating AIDS)
       144114-21-6, Retropepsin
RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
IT
              (prepn. of sulfonylamino acid amides contg. cis epoxide as irreversible
        HIV protease inhibitors for treating AIDS) 74-88-4, Iodomethane,
TΤ
                             78-77-3, Isobutyl bromide 78-82-0, Isobutyronitrile
        88-14-2, 2-Furancarboxylic acid 96-41-3; Cyclopentyl alcohol
                                      98-59-9, Toluenesulfonyl chloride 98-98-6,
        2-Furanmethanol
                                                        100-39-0, Benzyl bromide, 100-46-9,
        2-Pyridinecarboxylic/acid
       Benzylamine, reactions: 100-55-0, 3-Pyridylcarbinol 110-68-9, Methyl-N-butylamine 501-53-1, Benzyl chloroformate 503-74-2, Isovaleric acid 527-72-0, 2-Thiophenic acid 574-98-1, N-(2-Bromoethyl)phthalimide 586-95-8, 4-Pyridylcarbinol 586-2-Pyridylcarbinol 586-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 150-2-1, 
                                                                                                              586-98-1,
                                        603-35-0, Triphenylphosphine, reactions
        2-Pyridylcarbinol
        2-Furanylmethylamine 625-45-6; Methoxyacetic acid
                                    1779-49-3, Methyltriphenylphosphonium bromide
        L-Penicillamine
        2516-33-8, Cyclopropylmethanol 2516-47-4, Cyclopropylmethylamine
        4083-57-2, 3-Amino-2, 4-dimethylpentane, 6921-34-2, Benzylmagnesium
                        7693-46-1, p-Nitrophenyl chloroformate 13734-34-4
        23844-66-8, (R)-1-Amino-2-methyl-1-phenylpropane 24424-99-5,
        Di-tert-Butyl dicarbonate 24939-24-0, p-Aminobenzenesulfonyl chloride 33445-07-7, Isopropoxyacetic acid 37222-66-5, Oxone 59830-60-3,
        N-Benzyloxycarbonyl-L-phenylalaninal 68906-26-3, (S)-1-Amino-2-methyl-1-phenylpropane 69492-74-6, Thiopheneacetic acid 74124-79-1,
        Disuccinimidyl carbonate 96928 87-9 111491-96-4
Furanacetic acid 174562-82-4
RL: RCT (Reactant)
                                                                                                     137867-58-4,
              (prepn. of sulfonylamino acid amides contglicis epoxide as irreversible
             HIV protease inhibitors for treating AIDS)
                                                        97589-56-5P 100217-05-8P 112898-22-3P
IT
        65273-64-5P
                                82894-53-9P
                                  156641-80-4P [156641-81-5P] 156641-82-6P
                                                                                                              156641-83-7P
        156641-79-1P
                                                           160742+45-0P 174562-83-5P 174562-84-6P
                                  160742-44-9P
        156715-06-9P
                                                           174562-87-9P 3 174562 88 0P 1174562-89-1P
        174562-85-7P
                                 174562-86-8P
                                                        174562-92#6P|| 174562-93-7P|
        174562-90-4P 174562-91-5P
                                                           174689-87-3P
                                 174689'-86-2P
        174562-94-8P
        RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
              (prepn. of sulfonylamino acid amides contg. cis epoxide as irreversible
```

Absolute stereochemistry.

```
L27
                                     1995:994162 HCAPLUS 124:87790
                             .
AN
DN
     Pharmaceutical compositions containing HIV protease inhibitors and their preparation.
ΤI
     Al-Razzak, Laman; Marsh, Kennange.; Manning, Lourdes P.; Kaul, Dilip
IN
     Abbott Laboratories, USA
PCT Int. Appl., 58 pp.
CODEN: PIXXD2
Patent
English
PA
SO
DT
LΑ
     English
     ICM A61K031-425
IC
     ICS A61K009-08; A61K047-10; A61K047-12; A61K047-14
     34-2 (Amino Acids, Peptides, and
CC
     Proteins)
     Section cross-reference(s): 63
FAN.CNT 1
                                     Hay parties
                                      APPLICATION NO.
                                                                 DATE
     PATENT NO.
                       KIND DATE
                        A1 19950803. WO 1995-US232
                                                                 19950103
PΙ
     WO 9520384
         W: AU, CA, JP, KR, MX
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                        A1 20000726 TL 1994-111991 19941215
AA 19950803 CA 1995-2178632 19950103
     IL 111991
     CA 2178632
         19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103

19950103
     AU 9515248
     AU 700942
     EP 732923
                      T2 19970826 JP 1995-520059
     JP 09508383
                                                                19950103
                        A 19960116 US 1995-440277
A 19940128
                                                                 19950512
     US 5484801
PRAI US 1994-189021
                         A 19940729
     US 1994-283239
                           19950103,
     WO 1995-US232
                         W
GΙ
```

A pharmaceutical compn. which comprises a soln. of an HIV protease AB inhibiting compd. (e.g., I) in a pharmaceutically acceptable org. solvent comprising a mixt. of (1): (a) a solvent selected from propylene glycol and polyethylene glycol or (b) a solvent selected from polyoxyethyleneglycerol triricinoleate, polyethylene glycol 40 hydrogenated castor oil, fractioned coconut oil; polyoxyethylene 20 sorbitan monooleate and 2-(2-ethoxyethoxy) ethanol or (c) a mixt. thereof; and (2) ethanol or propylene glycol, is claimed. I was prepd. in many steps and its bioavailability in various formulations was studied. hiv protease inhibitor pharmaceutical compn; valinyldiphenylhexane prepn ST hiv protease inhibitor ΙT Pharmaceutical dosage forms (pharmaceutical compns. contg. HIV protease inhibitors and their prepn.) 155213-67-5P IT RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (pharmaceutical compns. contg. HIV protease inhibitors and their prepn.) IT 143838-10-2P 144164-10-3P RL: BYP (Byproduct); PREP (Preparation) (pharmaceutical compns. contg. HIV protease inhibitors and their prepn.)
62-56-6, Thiourea, reactions, 75-12-7, Formamide, reactions ΙT Ethyl chloroacetate: 109-94-4; Ethyl formate: 534-07-6; 1,3-Dichloroacetone: 563-83-7, Tsöbutyramide 3,4070-48-8, Valine methyl 40635-67-4, 6372-14-1, N-Benzyloxycarbonylphenylalaninol .alpha.-Acetoxyisobutyryl bromide 1156732 13-7 156732-15-9 ារីស ១៦ ខាង ប្រ RL: RCT (Reactant) (pharmaceutical compns. contg. HIV protease inhibitors and their prepn.) ΙT 13515-65-6P 32955-21-8P, 2-Amino-5-115-08-2P, Thioformamide 32955-22-9P, Ethyl thiazole-5-carboxylate (ethoxycarbonyl)thiazole 33142-21-1P, Ethyl 2-chloro-2-formylacetate 38585-74-9P, 5-Hydroxymethylthiazole 59830-60-3P 65386-28-9P 137649-69-5P 144163-44-0P 144163-85-9P#, 144163-97-3P 144164-11-4P 144141-68-4P \*154212-61-0P||| 154248-99-4P 154212-60-9P 154212-59-6P 162849-93-6P 162849-94-7P 162537-10-2P 162849-92-5P 162849-96-9P, 2-Amino-5-(ethoxycarbonyl)thiazole 162849-95-8P 165315-39-9P 162990-03-6P hvdrochloride RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (pharmaceutical compas. contg HIV protease inhibitors and their prepn.) IT 162990-01-4

红斑 精红 计

(pharmaceutical compns. contg HIV protease inhibitors and their

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

prepn.)

IT 162537-10-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (pharmaceutical compns. contg. HIV protease inhibitors and their prepn.)

RN 162537-10-2 HCAPLUS

CN L-Valine, N-[(4-nitrophenoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L27 ANSWER 13 OF 24 HCAPLUS COPYRIGHT 2001 ACS.

AN 1995:965043 HCAPLUS

DN 124:117909

Optimization of Alkylating Agent Prodrugs Derived from Phenol and Aniline Mustards: A New Clinical Candidate Prodrug (ZD2767) for Antibody-Directed Enzyme Prodrug Therapy

AU Springer, Caroline J.; Dowell, Robert; Burke, Philip J.; Hadley, Elma; Davies, D. Huw; Blakey, David C.; Melton, Roger G.; Niculescu-Duvaz, Ion

CS Cancer Research Campaign Centre for Cancer Therapeutics, Institute of Cancer Research, Sutton, SM2 5NG, UK

SO J. Med. Chem. (1995), 38(26), 5051-65 CODEN: JMCMAR; ISSN: 0022-2623

DT Journal

LA English

CC 34-2 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 1

GΙ

AB Sixteen novel potential prodrugs I [R = H, 2-Me, 2-Cl, 3-Me, 3-Me2CH, 3-F, 2,3-(CH:CHCH:CH), 3-CN; Z = O, NH; X, Y = Cl, Br, iodo, O3SMe] derived from phenol or aniline mustards and their 16 corresponding drugs II with ring substitution and/or different alkylating functionalities were designed. They are bifunctional alkylating agents in which the activating effect of the phenolic hydroxyl or amino function is masked through an oxycarbonyl or a carbamoyl bond to a glutamic acid. These prodrugs were

designed to be activated to their corresponding phenol and aniline nitrogen mustard drugs at a tumor site by prior administration of a monoclonal antibody conjugated to the bacterial enzyme carboxypeptidase G2 (CPG2) in antibody-directed enzyme prodrug therapy (ADEPT). The synthesis of the analogous novel parent drugs II is also described. The viability of a colorectal cell line (LoVo) was monitored with the potential prodrugs and the parent drugs. The differential in the cytotoxicity between the potential prodrugs and their corresponding active drugs ranged between 12 and >195 fold. Some compds. I exhibited substantial prodrug activity, since a cytotoxicity differential of >100 was achieved compared to the analogous II. The ability of the potential prodrugs to act as substrates for CPG2 was detd. (kinetic parameters KM and kcat), and the chem. stability was measured for all the compds. The unsubstituted phenols with different alkylating functionalities (I; R = H, Z = O) proved to have the highest ratio of substrates kcat:KM. From these studies, III (ZD2767) emerges as a new ADEPT clin. trial candidate due to its physicochem. and biol. characteristics.

ZD 2767 prepn biol alkylating agent; glutamate aniline mustard prepn ST ADEPT; phenol glutamate mustard prepn ADEPT; structure activity alkylating agent prodrug

ΙT Neoplasm inhibitors

(optimization of alkylating agent prodrugs derived from phenol and aniline mustards in prepn. of prodrug ZD2767 for antibody-directed enzyme prodrug therapy)
Alkylating agents, biological

ΙT

RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(optimization of alkylating agent prodrugs derived from phenol and aniline mustards in prepn. of prodrug ZD2767 for antibody-directed A THE PARTY OF THE enzyme prodrug therapy)

अभी है के इंकर्स है के अपने पूर्व ।

IT Antibodies

Enzymes

Apple of the Control RL: BPR (Biological process); BIOL! (Biological study); PROC (Process) (optimization of alkylating agent prodrugs derived from phenol and aniline mustards in prepn. of prodrug ZD2767 for antibody-directed and the first enzyme prodrug therapy)

Molecular structure-biological activity relationship ΙT (cytotoxic, optimization of alkylating agent prodrugs derived from phenol and aniline mustards in preph. of prodrug ZD2767 for antibody-directed enzyme prodrug therapy)

ΙT Pharmaceutical dosage forms

> (prodrugs, optimization of alkylating agent prodrugs derived from phenol and aniline mustards in prepn. of prodrug ZD2767 for antibody-directed enzyme prodrug therapy)

156079-29-7P 156079-30-0P 156079-35-5P 172974-00-4P 156079-29-7P 156079-31-1P IT 21667-05-0P 156078-84-1P 156079-32-2P 156079-34-4P 172974-01-5P 172974-03-7P 172974-19-5P 172974-20-8P 172974-21-9P 172974-02-6P 172974-22-0P

RL: BAC (Biological activity or effector, except adverse); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (optimization of alkylating agent prodrugs derived from phenol and aniline mustards in prepn. of prodrug ZD2767 for antibody-directed enzyme prodrug therapy)

1156078-94-3P 156078-98-7P 156079-02-6P 156078-91-0P IT 156078-82-9P 156079-04-8P 156079-05-9P 156079-06-0P 156079-07-1P 156079-03-7P 156079-88-8P, ZD 2767 156079-91-3P 156079~56-0P 156079-57-1P 172974~23-1P 172974-24-2P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (optimization of alkylating agent prodrugs derived from phenol and

```
aniline mustards in prepn. of prodrug ZD2767 for antibody-directed
                                      enzyme prodrug therapy)
    75-21-8, Oxirane, reactions 111-42-2, reactions 350-30-1, 3-Chloro-4-fluoronitrobenzene 399-96-2, 4-Amino-2-fluoropheno 2791-84-6 2834-90-4,
ΙT
                                                             2835-99-6,
     4-Amino-1-naphthol 2835-96-3, 4-Amino-2-methylphenol
                                                                   5854-73-9
     4-Amino-3-methylphenol 3964-52-1, 4-Amino-2-chlorophenol
     7693-46-1, 4-Nitrophenyl chloroformate 17417-09+3, 2-Fluoro-5-
                         17609-80-2, 4-Amino-3-chlorophenol 32677-01-3,
     nitrobenzonitrile
     Di-tert-butyl glutamate hydrochloride 82774-61-6 156639-14-4
     172974-26-4
     RL: RCT (Reactant)
        (optimization of alkylating agent, prodrugs derived from phenol and
        aniline mustards in prepn. of prodrug 2D2767 for antibody-directed
        enzyme prodrug therapy)
                                                156078-85-2P
                                                                156078-86-3P
ΙT
     65976-57-0P
                   65976-66-1P
                                 156078-83-0P
                    156078-93-2P
                                   156078-95-4P
                                                  156078-99-8P
     156078-92-1P
                    156079-01-5P
                                   156079-08-2P
                                                  156079-09-3P
     156079-00-4P
                                   156079-12-8P
                                                  156079-13-9P
                                                                  156079-15-1P
     156079-10-6P
                    156079-11-7P
                                   156079-18-4P 156079-19-5P
                                                                  156079-20-8P
     156079-16-2P
                    156079-17-3P
                                                  156079-25-3P
                                                                  156079-27-5P
                    156079-23-1P
                                   156079-24-2P
     156079-22-0P
                                   156079-37-7P 3 156079-38-8P;
                                                                  156079-39-9P
                    156079-36-6P
     156079-28-6P
                                                  :156079-59-3P
                    156079-41-3P
                                   156079-58-2P
                                                                  156079-60-6P
     156079-40-2P
                                   156079-66-2P+: 156079-89-9P+
     156079-61-7P
                    156079-65-1P
                                                                 156079-90-2P
                    156639-24-6P 156639-26-8P 172973-90-9P 172973-93-2P 172973-94-3P 172973-95-4P
                                                                 172973-91-0P
     156079-92-4P
                                                                  172973-96-5P
     172973-92-1P
                                   172973-99-8PF 172974-04-8P
                    172973#98-7P
                                                                 ·172974-05-9P
     172973-97-6P
                                   172974-08-2P 172974-09-3P
                                                                  172974-10-6P
     172974-06-0P
                    172974÷07-1P
                                   172974-13-9P # 172974-14-0P
                                                                 172974-15-1P
                    172974+12-8P
     172974-11-7P
                                   172974-18-4P 11172974-25-3P
                    172974-17-3P
     172974-16-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (optimization of alkylating agent prodrugs derived from phenol and
        aniline mustards in prepn. of prodrug ZD2767 for antibody-directed
                                   enzyme prodrug therapy)
ΙT
     156079-00-4P 172974-16-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (optimization of alkylating agent prodrugs derived from phenol and
        aniline mustards in prepn. of prodrug ZD2767 for antibody-directed
                                   通行机 计
        enzyme prodrug therapy)
                                      中间 同篇 出。
RN
     156079-00-4 HCAPLUS
     L-Glutamic acid, N-[(4-nitrophenoxy)carbonyl]-, bis (1,1-dimethylethyl)
CN
                  (CA INDEX NAME)
     ester (9CI)
                                    Laidylake gr
Absolute stereochemistry.
                                   1274年1186年11
                                     OP:
                                    OBú-ting in i
                                    清晰神神神
                                   OBu-t
02N
                                   at Agent is at the lea
```

( II

RN 172974-16-2 HCAPLUS

CN L-Glutamic acid, N-[%3-fluoro-4-nitrophenoxy) carbonyl]-bis(1,1-dimethylethyl) ester (9CI) \* (CA INDEX NAME)

Absolute stereochemistry.

朝的激化

```
F O H N S OBu-t
```

```
COPYRIGHT 2001 ACS
L27
     ANSWER 14 OF 24 HCAPLUS
     1995:958521 HCAPLUS
AN
DN
     124:176946
     Preparation of retroviral protease inhibiting peptide analogs.
ΤI
     Norbeck, Daniel W.; Sham, Hing L.; Kempf, Dale J.; Zhao, Chen
IN
     Abbott Laboratories, USA
PA
     U.S., 66 pp. Cont.-in-part of U.S. Ser, No. 23,226, abandoned.
so
     CODEN: USXXAM
                          1 :
                                     经建筑 海豚
DT
     Patent
                                    2-14;
LΑ
     English
IC
     ICM A61K031-44
          A61K031-425; A61K031-42; C07D413-14; C07D417-14; C07D263-30;
          C07D277-20
NCL
     514333000
     34-3 (Amino Acids, Peptides, and and
CC
                                     1.16
     Section cross-reference(s): 1
FAN.CNT 2
                                            APPLICATION NO.
                       KIND DATE
     PATENT NO.
                                        US 1994-185666
                                                               19940201
                             19951024
     US 5461067
PΙ
                        A1 19940901 WO 1994-US1457
                                                               19940208
     WO 9419332
             CA, JP
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
155338 AA 199409018 CA 1994-2155338 19940208
     CA 2155338
                        A1 :: 19951129 Ep. 1994-908018
                                                               19940208
     EP 683772
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
                        T2
                            19960730 *** *** JP | 1994-519025
                                                               19940208
     JP 08507061
                        A 19970415
                                            US 1995-455922
                                                               19950531
     US 5621109
                            19970520
                                        9-45 EUS 1995+455458
                                                               19950531
     US 5631376
                            19950531
     US 5990135
                             19930225
PRAI US 1993-23226
                          19940201
19940208
     US 1994-185666
     WO 1994-US1457
os
     MARPAT 124:176946
     R6YmNHCHR1CH(OH)CH2NR2NH(Y1)nR5: [R1, R2 = H, alkyl, aryl, thioalkoxyalkyl,
AB
     aralkyl, cycloalkyl, guanidinoalkyl, arylthioalkoxyalkyl,
     cycloalkyloxyalkyl, cycloalkylsulfonylalkyl, aminocarbonylalkyl, etc.; Y =
     NHCHR4CO, NHNR4CO, etc.; Y1 = COCHR3NH, CONR3NH, etc.; R5, R6 = C(:T)GR7;
     T = O, S; G = CH2, O, S, NR8; R8 = H, alkyl, cycloalkyl; R7 = alkyl,
     cycloalkyl, aryl, arylalkyl, arylalkoxyalkyl, aminoalkyl, N-protected
     aminoalkyl, alkylaminoalkyl, N-protected alkylaminoalkyl,
     dialkylaminoalkyl, carboxyalkoxyalkyl, (alkoxycarbonyl)alkoxyalkyl,
     carboxyalkyl, alkoxycarbonylalkyl, aminocarboxyalkyl, N-protected
     aminocarboxyalkyl, alkylaminocarboxyalkyl, etc.; m, n = 0, 1], were prepd.
     Thus, (5S)-[[(5-thiazolyl)methoxy]carbonyl]amino-2-[(N-2-thiazolyl)methoxy]carbonyl]amino-2-[(N-2-thiazolyl)methoxy]
     oxazoly1)methoxycarbony1]amino-45-hydroxy-1-(3-furany1)-6-pheny1-2-
     azahexane (soln. phase prepn. given) inhibited HIV-13B in MT4 cells with IC50 = 0.029-0.032 .mu.M.
     peptide analog prepn retroviral protease inhibitor; aminopropylhydrazine
ST
```

11.

李克斯蒙蒙亚亚 (1975年)

1917 1

```
azolylmethoxycarbonyl amino acid prepn virucide
IT
     Virucides and Virustats
        (prepn. of retroviral protease inhibiting peptide analogs)
ΙT
     Peptides, preparation;
     RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation): USES: (Uses)
     (Preparation); USES: (Uses)
        (analogs, prepn. of retroviral protease inhibiting peptide analogs)
ΙT
     Virus, animal
        (human immunodeficiency 1, treatment of HIV infections; prepn. of
        retroviral protease inhibiting peptide analogs)
ΙT
     144114-21-6, Retropepsin
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (HIV protease inhibitors; prepn. of retroviral protease inhibiting
        peptide analogs)
                     150767-07-0P
                                     162739-20-0P 162739-21-1P
                                                                    162739-22-2P
ΙT
     150767-06-9P
                     162739-24-4P
                                    162739-25-5P
                                                    162739-26-6P
                                                                    162739-27-7P
     162739-23-3P
                    162739-31-3P
                                    162739-32-4P 162739-33-5P
                                                                    162739-38-0P
     162739-28-8P
                                    162739-41-5P
                                                  ; 162739-42-6P
                                                                    162739-43-7P
     162739-39-1P
                     162739÷40-4P
                                                    173767-18-5P
                                                                    173767-19-6P
     162739-46-0P
                     173767-16-3P
                                    173767-17-4P
                     173767-21-0P
                                     173767-22-1P
                                                    173767-23-2P
                                                                    173767-24-3P
     173767-20-9P
     173767-25-4P
                    173767-26-5P
                                    173767-27-6P 173767-28-7P
                                                                    173767-29-8P
     173767-30-1P
                    173767-31-2P
                                    173767-32-3P
                                                   173767-33-4P
                                                                    173767-34-5P
                                   173767-37-8P
                                                    173767-38-9P
173767-43-6P
                                                                   :173767-39-0P
     173767-35-6P
                     173767-36-7P
                     173767-41-4P
                                     173767-42-5P
                                                                    173767-44-7P
     173767-40-3P
                                   173767-47-0P 173767-48-1P
173767-52-7P 173767-53-8P
                     173767-46-9P
     173767-45-8P
                                                                    173767-49-2P
                     173767-51-6P
                                                                    173767-54-9P
     173767-50-5P
                                                    173767-58-3P
                                     173767-57-2P
                                                                    173767-59-4P
                     173767-56-1P
     173767-55-0P
                    173767-61-8P 173767-62-9P 173767-63-0P 173767-66-3P 173767-67-4P 173767-68-5P
                                                                    173767-64-1P
     173767-60-7P
                                                                    173767-69-6P
     173767-65-2P
                                     173767-72-1P | 173767-73-2P
                                                                    173767-74-3P
     173767-70-9P
                     173767-71-0P
                                    173767-77-6P 173767-78-7P
     173767-75-4P
                     173767-76-5P
                                                                    173767-79-8P
                     173767-81-2P
                                     173767-82-3P
                                                    173767-83-4P
                                                                    173767-84-5P
     173767-80-1P
                                     173767-87-8P 173767-88-9P
     173767-85-6P
                     173767-86-7P
                                                                    173767-89-0P
                                   173767-92-5P 173767-93-6P
                                                                   173767-94-7P
     173767-90-3P
                     173767-91-4P
                                    173767-97-0P
                                                    173767-98-1P
                                                                    173767-99-2P
     173767-95-8P
                     173767-96-9P
                                    173768-02-0P
                                                                    173768-04-2P
                     173768-01-9P
                                                    173768-03-1P
     173768-00-8P
                     173768-06-4P
                                     173768-07-5P
                                                    173768-08-6P
                                                                    173768-09-7P
     173768-05-3P
                     173768-11-1P
                                    173768-12-2P
                                                   : 173768-13-3P
                                                                    173768-14-4P
     173768-10-0P
                                     173768-17-7P
                                                    173768-18-8P
                                                                    173768-19-9P
     173768-15-5P
                    .173768-16-6P
                                    173768-22-4P
                                                                    173768-24-6P
                     173768-21-3P
                                                    173768-23-5P
     173768-20-2P
                                                                    173768-29-1P
     173768-25-7P
                     173768-26-8P
                                     173768-27-9P
                                                    173768-28-0P
                                                   173768-33-7P
                                                                    173768-34-8P
     173768-30-4P
                     173768-31-5P
                                     173768-32-6P
                                                    173768-38-2P
                                                                    173768-39-3P
     173768-35-9P
                     173768-36-0P
                                     173768-37-1P
                     173768-41-7P
                                    173768-42-8P
                                                    173768-43-9P
                                                                    173768-44-0P
     173768-40-6P
                                     173768-47-3P
                                                    173768-48-4P
                                                                    173768-49-5P
     173768-45-1P
                     173768-46-2P
     173768-50-8P
                     173768-51-9P
                                     173768÷52-0P
                                                    173768+53-1P
                                                                    173768-54-2P
                                   173768-57-5P
                                                   173768+58-6P
                                                                    173768-59-7P
     173768-55-3P
                     173768-56-4P
     173768-60-0P
                     173768-61-1P
                                    173768-62-2P
                                                   . 173768+63-3P
                                                                    173768-64-4P
     173768-65-5P
                     173768-66-6P
                                    173768-67-7P
                                                   173768+68-8P
                                                                    173768-69-9P
                                                                    173768-74-6P
     173768-70-2P
                     173768-71-3P
                                     173768-72-4P
                                                    173768+73-5P
                                     173768-77-9P 173768-78-0P
                                                                    173768-79-1P
     173768-75-7P
                     173768-76-8P
                                    173768-82-6P
                                                                    173768-84-8P
                                                    173768-83-7P
     173768-80-4P
                     173768-81-5P
                                                    173768-88-2P
                                                                    173768-89-3P
                     173768-86-0P
                                     173768-87-1P
     173768-85-9P
                                     173768-92-8P 173768-93-9P
                                                                    173768-94-0P
                     173768-91-7P
     173768-90-6P
                                     173768-97-3P 173768-98-4P
                                                                    173768-99-5P
     173768-95-1P
                     173768-96-2P
                     173769-01-2P
                                     173769-02-3P
                                                    173769-03-4P
                                                                    173769-04-5P
     173769-00-1P
                     173769-06-7P
                                    173769-07-8P
                                                    173769-08-9P
                                                                    173769-09-0P
     173769-05-6P
                                     173769-12-5P
                                                                    173769-14-7P
                     173769-11-4P
                                                    173769-13-6P
     173769-10-3P
                     173769-16-9P
                                     173769-17-0P
                                                    173769-18-1P
                                                                    173769-19-2P
     173769-15-8P
```

,秦军的范围的"静静""诸侯",

经济制度

```
Total design of the second sec
                                                       173769-22-7P
                                                                               173769-23-8P
       173769-20-5P
                               173769-21-6P
                                                                                                       173769-24-9P
                               173769-26-1P
                                                       173769-27-2P
                                                                               173769-28-3P
                                                                                                       173769-29-4P
       173769-25-0P
       RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic
       preparation); THU (Therapeutic use); BIOL (Biological study); PREP
        (Preparation); USES' (Uses)
             (prepn. of retroviral protease inhibiting peptide analogs)
                                                       173769-32-9P
                                                                               173769-33-0P
                                                                                                       173769-34-1P
IT
                               173769-31-8P
       173769-30-7P
                                                       173769-37-4P
                                                                               173769-38-5P
                                                                                                       173769-39-6P
       173769-35-2P
                               173769-36-3P
                                                       173769-42-1P
                                                                               173769-43-2P
                                                                                                       173769-44-3P
       173769-40-9P
                               173769-41-0P
                                                                               173769-48-7P
                                                                                                       173769-49-8P
       173769-45-4P
                               173769-46-5P
                                                       17,3769-47-6P
                                                                                                       173769-54-5P
                                173769-51-2P
                                                       173769-52-3P
                                                                               173769-53-4P
       173769-50-1P
                               173769-56-7P
                                                       173769-57-8P
                                                                               173769-58-9P
                                                                                                       173769-59-0P
       173769-55-6P
                                                       173769-62-5P
                                                                               173769-63-6P
                                                                                                       173769-64-7P
       173769-60-3P
                               173769-61-4P
                                                       173769-67-0P
                                                                               173769-68-1P
                                                                                                       173769-69-2P
                               173769-66-9P
       173769-65-8P
       173769-70-5P
                               173769-71-6P
                                                       173769-72-7P
                                                                               173769-73-8P
                                                                                                       173769-74-9P
                               173769-76-1P
       173769-75-0P
                                                       173769-77-2P
                                                                               173769-78-3P
                                                                                                       173769-79-4P
                               173769-81-8P
                                                       173769-82-9P
                                                                               173769-83-0P
                                                                                                       173769-84-1P
       173769-80-7P
                               173769-86-3P
                                                       17,3769-87-4P
                                                                               173769-88-5P
                                                                                                       173769-89-6P
       173769-85-2P
                                                       173769-92-1P
                                                                            173769-93-2P
                                                                                                       173769-94-3P
                               173769-91-0P
       173769-90-9P
                                                       173769-97-6P
                                                                               173769-98-7P
                               173769-96-5P
                                                                                                       173769-99-8P
       173769-95-4P
                                                       173770-02-0P
                                                                               173770-03-1P
                                                                                                       173770-04-2P
       173770-00-8P
                               173770-01-9P
                                                        173770-07-5P
                                                                               173770-08-6P
                                                                                                       173770-09-7P
       173770-05-3P
                                173770-06-4P
                                                       173770-12-2P
                                                                               173770-13-3P
                                                                                                       173770-14-4P
       173770-10-0P
                                173770-11-1P
                                173770-16-6P
                                                      173770-17-7P 173770-18-8P
                                                                                                       173770-19-9P
       173770-15-5P
                                                       173770-22-4P (173770-23-5P)
       173770-20-2P
                               173770-21-3P
                                                                                                       173770-24-6P
                                                                               173770-28-0P
                                                       173770-27-9P
                                                                                                       173770-29-1P
       173770-25-7P
                               173770-26-8P
                               173770-31-5P
                                                      173770-32-6P
                                                                               173770-33-7P
                                                                                                       173770-34-8P
       173770-30-4P
                                                                               173770-38-2P
                                                       173770-37-1P
                                                                                                       173770-39-3P
                               173770-36-0P
       173770-35-9P
                                                       173770-42-8P
                                                                               173770-43-9P
                                                                                                       173770-44-0P
                               173770-41-7P
       173770-40-6P
                                                       173770-47-3P
                                                                               173770-48-4P
                                                                                                       173770-49-5P
                                173770-46-2P
       173770-45-1P
                                                       173770-52-0P
                                                                               173770-53-1P
                                                                                                       173770-54-2P
                               173770-51-9P
       173770-50-8P
                                                                                                       173770-59-7P
                                173770-56-4P
                                                       173770-57-5P
                                                                               173770-58-6P
       173770-55-3P
                                                                               173770-63-3P
                                                                                                       173770-64-4P
       173770-60-0P
                               173770-61-1P
                                                       173770-62-2P
                                                                                                       173770-69-9P
                                                       173770-67-7P
                                                                               173770-68-8P
       173770-65-5P
                                173770-66-6P
                                                                                                       173770-74-6P
                                                       173770-72-4P
                                                                               173770-73-5P
        173770-70-2P
                                173770-71-3P
                                                                                                       173770-79-1P
       173770-75-7P
                                173770-76-8P
                                                       173770-77-9P
                                                                               173770-78-0P
                                173770-81-5P
                                                       173770-82-6P
                                                                               173770-83-7P
                                                                                                       173770-84-8P
       173770-80-4P
                               173770-86-0P
                                                       173770-87-1P
                                                                               173770-88-2P
                                                                                                       173770-89-3P
       173770-85-9P
       173770-90-6P
                               173770-91-7P
                                                       173770-92-8P
                                                                               173770-93-9P
                                                                                                       173770-94-0P
                                                                               173770-98-4P
                                                                                                       173770-99-5P
       173770-95-1P
                                173770-96-2P
                                                       173770-97-3P
                                                       173771-02-3P
                                                                               173771-03-4P
                                                                                                       173771-04-5P
       173771-00-1P
                               173771-01-2P
                                173771-06-7P
                                                       173771-07-8P
                                                                               173771-08-9P
                                                                                                       173771-09-0P
       173771-05-6P
                                                                               173771-13-6P
                                                       173771-12-5P
                                                                                                       173771-14-7P
                                173771-11-4P
       173771-10-3P
                                                       173771-17-0P
                                                                               173771-18-1P
                                173771-16-9P
                                                                                                       173771-19-2P
       173771-15-8P
                                                       173771-22-7P
                                                                              173771-23-8P
                                                                                                       173771-24-9P
        173771-20-5P
                                173771-21-6P
                                                       173771-27-2P 173771-28-3P
                                173771-26-1P
        173771-25-0P
                                                                                                       173771-29-4P
                                                        173771-32-9P
                                                                               173771-33-0P
                                                                                                       173771-34-1P
        173771-30-7P
                                173771-31-8P
                                                       173771-37-4P
                                                                               173771-38-5P
                                173771-36-3P
                                                                                                       173771-39-6P
        173771-35-2P
        173771-40-9P
                                173771-41-0P
                                                        173771-42-1P
                                                                               173771-43-2P
                                                                                                       173771-44-3P
                                                       173771-47-6P
                                                                               173771-48-7P
                                                                                                       173771-49-8P
        173771-45-4P
                                173771-46-5P
                                                                               173771-53-4P
                                                                                                       173771-54-5P
                                173771-51-2P
                                                       173771-52-3P
        173771-50-1P
        173771-55-6P
                                                       173771-57-8P
                                                                               173771-58-9P
                                                                                                       173771-59-0P
                                173771-56-7P
                                                                               173771-63-6P
                                                                                                       173771-64-7P
                                173771-61-4P
                                                       173771-62-5P
        173771-60-3P
       RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
       preparation); THU (Therapeutic use); BIOL (Biological study); PREP
        (Preparation); USES (Uses)
             (prepn. of retroviral protease inhibiting peptide analogs)
                                                                                                       173771-69-2P
IT
        173771-65-8P
                                173771-66-9P
                                                        173771-67-0P
                                                                               173771-68-1P
                                173771-71-6P
                                                        173771-72-7P
                                                                               173771-73-8P
                                                                                                       173771-74-9P
        173771-70-5P
                                                       173771-77-2P
                                173771-76-1P
                                                                               173771-78-3P
                                                                                                       173771-79-4P
        173771-75-0P
                                       1 (i
                                                       到11111
```

n | 

A PASTY

41 1

```
173771-80-7P
               173771-81-8P
                               173771-82-9P
                                               173771-83-0P
                                                               173771-84-1P
                                              173771-88-5P
                               173771-87-4P
                                                               173771-89-6P
173771-85-2P
               173771-86-3P
                                               173771-93-2P
                               173771-92-1P
                                                               173771-94-3P
173771-90-9P
               173771-91-0P
                                                               173771-99-8P
               173771-96-5P
                               173771-97-6P
                                               173771-98-7P
173771-95-4P
               173772-01-5P
                               173772-02-6P
                                               173772-03-7P
                                                               173772-04-8P
173772-00-4P
                               173772-07-1P
                                               173772-08-2P
                                                               173772-09-3P
               173772-06-0P
173772-05-9P
                               173772-12-8P
                                               173772-13-9P
                                                               173772-14-0P
               173772-11-7P
173772-10-6P
                                                               173772-19-5P
                                               173772-18-4P
               173772-16-2P
                               173772-17-3P
173772-15-1P
                               173772-22-0P
                                               173772-23-1P
                                                               173772-24-2P
173772-20-8P
               173772-21-9P
                               173772-27-5P
                                               173772-28-6P
                                                               173772-29-7P
173772-25-3P
               173772-26-4P
                                               173772-33-3P
                                                               173772-34-4P
               173772-31-1P
                               173772-32-2P
173772-30-0P
               173772-36-6P
                               173772-37-7P
                                               173772-38-8P
                                                               173772-39-9P
173772-35-5P
                                               173772-43-5P
                               173772-42-4P
                                                               173772-44-6P
               173772-41-3P
173772-40-2P
               173772-46-8P
                               173772-47-9P
                                               173772-50-4P
173772-45-7P
RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); USES (Uses)
   (prepn. of retroviral protease inhibiting peptide analogs)
                               70-23-5, Ethyl bromopyruvate
                                                                74-89-5.
67-64-1, Acetone, reactions
                         75-12-7; Formamide, reactions 75-44-5, Carbonic
Methanamine, reactions
                                               93-55-0, Propiophenone
             89-98-5, 2-Chlorobenzaldehyde
dichloride
98-01-1, 2-Furaldehyde, reactions 98-83-9, reactions 98-86-2, Acetophenone, reactions 100-52-7, Benzaldehyde, reactions 100
Pyridine-3-methanol 100-83-4 Hydroxybenzaldehyde 104-87-0,
                 104-88-1, 4-Chlorobenzaldehyde, reactions
p-Tolualdehyde
                                                                105-39-5.
Ethyl chloroacetate 109-94-4 Ethyl formate 110-91-8, Morpholine, reactions 122-78-1, Phenylacetaldehyde 123-08-0, 4-Hydroxybenzaldehyde
123-11-5, p-Anisaldehyde, reactions 123-75-1, Pyrrolidine, reactions
459-57-4, 4-Fluorobenzaldehyde
                                                             529-20-4,
                 534-07-6, 1,3 Dichloroacetone 563-83-7, Isobutyramide
o-Tolualdehyde
586-98-1, Pyridine-2_methanol 587-04-2, 3-Chlorobenzaldehyde
                                                                    591-31-1
688-99-3, 3-Hydroxy-5-hexene 870-46-2, tert-Buty carbazate
                                                                   872-85-5,
                             925-90-6, Ethylmagnesium bromide
                                                                  930-45-0,
Pyridine-4-carboxaldehyde
                              1121-60-4 Pyridine-2-carboxaldehyde
(S,S)-2-Aminocyclopentanol
1779-49-3, Triphenylmethylphosphonium bromide 2043-61-0,
                                                                  6089-04-9
                             3731+51-9 2-Aminomethylpyridine
Cyclohexanecarboxaldehyde
6306-52-1, Valine methyl ester hydrochloride 6372-14-1 6972-05-0,
N, N-Dimethylthiourea 10200-59-6, 2-Thiazolecarboxaldehyde
                                                                 14337-43-0,
Ethyl chlorooximidoacetate
                              16332-06-2, 2-Methoxyacetamide
                                                                 74111-21-0,
                             82625-45-45 162740-04-7
                                                          162740-05-8
(S,S)-2-Aminocyclohexanol
                                173772-49-1
RL: RCT (Reactant)
   (prepn. of retroviral protease inhibiting peptide analogs)
                               934-53-2P 4-13242-92-7P
115-08-2P, Methanethioamide
                                                         13515-65-6P
14294-10-1P, 4-Morpholinecarbothioamide 15536 75 1P
                                                        16689-34-2P
                                           32939<del>-</del>32-5P
                                                         32955-22-9P
              24469-50-9P
                             30293+86-8P
16689~35-3P
                               hiazolemethanol 39624-97-0P 40398-3041337-78-4P 53370-84-6P 57699-48-6P
33142-21-1P
               38585+74-9P, 5-Thiazolemethanol
                                                                 40398-36-5P,
1-Pyrrolidinecarbothicamide
                             59830+60-3P 465386+28#9P 169353-16-8P
57699-55-5P
              57699-57-7P
98019-60-4P, 5-Isoxazolemethanol 99805-29-5P 100868-72-2P
                              130782-46-69 134807-06-0P
134807-30-0P 135941-95-6P
                                                               134807-20-8P
               126533-96-8P
126533-95-7P
                                                               135941-97-8P
134807-28-6P
               134807+29-7P
                               144163-71-3P 1144163-72-4P
                                                               144163-97-3P
               144163-45-1P
137515-66-3P
                                144186-59-4P 149267-56-1P
                                                               149267-64-1P
               144186-54-9P
144186-53-8P
                              150767-02-5P: 150767-04-7P
                                                               150767-05-8P
               149267-73-2P
149267-65-2P
                                150767-10-5P 154212-60+9P
                                                               154212-61-0P
150767-08-1P
                150767-09-2P
                               155884-24-5P 161852-61-5P
                154972-22-2P
154248-99-4P
                               162739-35-7P 1 162739-36-8P
162537-10-2P
              162739-34-6P
162739-37-9P
                                162739-48-2P 162739-49-3P 162739-50-6P
               162739-44-8P
                               162739-54-0P 162739-55-1P
                                                               162739-56-2P
162739-51-7P
                162739-52-8P
```

IT

```
经特别的 制度
                                     162739-59-5P
                                                    162739-60-8P
     162739-57-3P
                    162739-58-4P
                                                                    162739-61-9P
                                    162739-64-2P
                    162739-63-1P
                                                    162739-65-3P
                                                                    162739-66-4P
     162739-62-0P
                                     162739-69-7P | 162739-70-0P
                                                                    162739-71-1P
     162739-67-5P
                   162739-68-6P
                                    162739-74-4P
162739-81-3P
                                                    162739-75-5P
                                                                    162739-77-7P
     162739-72-2P 162739 \pm 73-3P
                                                  162739 82 4P 162739-83-5P
     162739-78-8P 162739-80-2P
                                    162739-86-8P 162739-88-0P
     162739-84-6P 162739-85-7P
                                                                    162739-89-1P
                                                    162739-93-7P
     162739-90-4P 162739-91-5P
                                    162739-92-6P
                                                                    162739-94-8P
                                    162739-97-1P
                                                    162739-98-2P
                                                                    162739-99-3P
     162739-95-9P
                    162739÷96-0P
                                    162740-02-5P
                                                  162740-03-6P
                                                                   163658-33-1P
     162740-00-3P
                    162740-01-4P
     173772-48-0P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of retroviral protease inhibiting peptide analogs)
     162537-10-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of retroviral protease inhibiting peptide analogs)
     162537-10-2 HCAPLUS
     L-Valine, N-[(4-nitrophenoxy)carbonyl], methyl ester (9CI) (CA INDEX
     NAME)
                                   Absolute stereochemistry.
                            OMe
    ANSWER 15 OF 24 HCAPLUS COPYRIGHT 2001 ACS 1995:695866 HCAPLUS
L27
     1995:695866 HCAPLUS
123:84005
     Preparation of peptide analogs as retroviral protease inhibitors.
Kempf, Dale J.; Norbeck, Daniel W.; Sham, Hing Leung; Zhao, Chen; Sowin,
     Thomas J.; Reno, Daniel S.; Haight Anthony R.; Cooper, Arthur J.
     Abbott Laboratories, USA
                                    Maria Maria
     PCT Int. Appl., 194 pp
     CODEN: PIXXD2
     Patent
                                    English
     ICM A61K031-425
     ICS A61K031-42; C07D277-24; C07D275-02; C07D261\pm08; C07D263-32
     34-3 (Amino Acids, Peptides, and
                                     Proteins)
     Section cross-reference(s): 1
FAN.CNT 6
                       KIND DATE APPLICATION NO. DATE

A1 19940707 WO 1993-US12326 19931216

JP, KR
     PATENT NO.
     WO 9414436
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
2170020 AA 19940630 CA 1993-2170020 19931216
2135890 AA 19940707 CA 1993-2135890 19931216
         W: AU, CA, JP, KR
     CA 2170020
                     AA 19940707 CA 1993-2135890 19931216
C 19960827;
A1 19940719 AU 1994-59546 19931216
     CA 2135890
     CA 2135890
     AU 9459546
                        B2 19950518
     AU 659575
```

B1 19960925

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE

A1 19951004

Male !

EP 1994-905429

19931216

EP 674513

EP 674513

IT

RN

CN

 $o_2N$ 

ANDN

ΤI IN

PA

SO

DT

LΑ

IC

CC

PI

```
T2
    JP 08505844
                            19960625
                                                           19931216
    JP 2637847
                      B2 19970806
                         19960821
                                          EP 1996-106301
                                                            19931216
    EP 727419
                      A2
    EP 727419
                            19961030
                      А3
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
                         , 1996101<u>5</u>
                                          AT 1994+905429
                                                            19931216
    AT 143262
                      \mathbf{E}
                      T3 | 19970201
                                          ES 1994-905429
                                                            19931216
    ES 2088839
                      A2 | 19970506
                                           JP 1996-132368
                                                            19931216
    JP 09118679
    JP 10087639
                      A2
                           19980407
                                           JP 1996-132369
                                                            19931216
                      A2 20010411
                                          EP 2000-124382
                                                            19931216
    EP 1090914
                            20010418
    EP 1090914
                      Α3
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE
                      A1 19950330 IL 1993-108126
A1 19950615 AU 1995-14927
                                                            19931221
    IL 108126
    AU 9514927
                                                            19950320
                      B2 19970424
    AU 677500
                           19960723
                      A'
                                           US 1995-410996
                                                            19950327
    US 5539122
                           19960903
                                          US 1995-411032
                                                            19950327
    US 5552558
                      Α
                      A
                                          US:1995-411140
    US 5696270
                            19971209
                                                            19950327
                                          ∵ÜS '1995-412253
                                                            19950328
    US 5580984
                      Α
                            19961203
                                          us 1995-412244
                                                            19950328
    US 5679797
                      A
                           19971021
    US 5583232
                            19961210
                                          US 1995-412821
                                                            19950329
                      Α
                                          US 1995-412438
    US 5597927
                      A:
                           19970128
                                                           . 19950329
                                    us 1995-413136
                            19971007
    US 5674882
                      A.
                                                            19950329
                                           US 1995-413290
                                                           19950330
    US 5583233
                      Α
                            19961210
    US 5625072
                      A 19970425
A 19970107
                                          US 1995-415827
                                                            19950403
                                         US 1995-416272
    US 5591860
                                                           19950404
                           19970128
                                         us 1995-416607
                      Α
                                                            19950404
    US 5597928
                      \mathbf{A}_{1}^{\dagger}
                            19970304
                                           US 1995-416259
                                                            19950404
    US 5608072
                      A 19901
A 19970819
                           19961015
                                                            19950405
                                           US 1995-417304
    US 5565418
                                           US 1995-417165
                                                            19950405
    US 5659044
                         19970819
                                           US 1995-417295
                                                            19950405
    US 5659045
                      Α
                            19970401
                                           US 1995-418056
                                                            19950406
                      Α
    US 5616720
                      A 19970603
                                         US 1995-417879
                                                           , 19950406
    US 5635523
                         19990406
    US 5892052
                      Α
                                          ∴ÿs 1995-418031
                                                            19950406
                           19960730
19981208
                                           ÜS 1995–423387
                                                            19950425
    US 5541206
                      A:
                                          US 1997-821609
                                                            19970320
    US 5846987.
                      A:
                            19990323
                                          ÜS 1997-822071
                                                            19970320
    US 5886036
                      Α
                      A1 19971211...
                                           AU 1997-28560
                                                            19970709
    AU 9728560
    AU 697681
                            1998101568
                                           B2
                            20000125
                                          US 1997-944351
                                                            19971006
    US 6017928
                      Α
                            20001121
                                          US 1998-207881
                                                            19981208
    US 6150530
                      Α
                            19921229
                                          3 P. A. 19
PRAI US 1992-998114
                      Α
                         19931202
                      A:
    US 1993-158587
       1983-355945
                       B2:
    US 1989-355945
                       B2:
                            19890523
                       B2 1
                            19890908
    US 1989-405604
                           19891222
    US 1989-456124
                       B2
                           19900509ht ....
    US 1990-518730
                      A2 .
                      B2 19901120
    US 1990-616170
    US 1991-746020
                       B2 19910815
                            19911023
                      B2
A3
    US 1991-777626
                            19931216
    CA 1993-2135890
                       А3
                            19931216
    EP 1994-905429
                       A3 19931216:
    EP 1996-106301
                      A3 19931216
     JP 1994-515323
                            19931216
    WO 1993-US12326
                       A3 19950329
    US 1995-413136
    US 1995-417879
                       A3 19950406
     US 1995-418031
                       A3 :
                            19950406
    MARPAT 123:84005
```

OS

AB Title compds. [I; R1 = monosubstituted thiazolyl, oxazolyl, isoxazolyl, isothiazolyl; n = 1-3; R2, R6 = H, alkyl; R3 = alkyl; R4, R41 = H(substituted) Ph, thiazolyl, oxazolyl; R7 = (alkyl-substituted) thiazolyl, oxazolyl, isoxazolyl, isothiazolyl; X = H, Y = OH, or X = OH, Y = H; Z =null, O, S, CH2, NR8; R8 = alkyl, cycloalkyl, OH; NH2, etc.; with provisos], were prepd. Thus, (2S, 3S, 5S) - 5 - [N - [N - [N - methyl - N - [(2 - methyl - [(2 - methyl - N - [(2 - methyl - N - [(2 - methyl - N - [(2 isopropyl-4-thiazolyl)methyl]amino]carbonyl]valinyl]amino]-2-[N-[(5thiazolyl)methoxycarbonyl]amino]-1,6-diphenyl-3-hydroxyhexane (prepn. via dimerization of Z-phenylalaninal given) inhibited HIV-13B in MT4 cells with IC50 = 0.025-0.040 .mu.M. peptide analog prepn hiv protease inhibitor; virucide heterocyclylpeptide ST 1 analog IT Virucides and Virustats (prepn. of peptide analogs as retroviral protease inhibitors) High the state of the Market 1 IT Peptides, preparation RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP . (Preparation); USES (Uses) (prepn. of peptide analogs as retroviral protease inhibitors) IT Virus, animal (human immunodeficiency, infection; prepn. of peptide analogs as retroviral protease inhibitors) 165314-94-3P 165314+95-4P 165314-93-2P 165314-96-5P ΙT 155213-67-5P 165314-99-8P 165315-00-4P 165315-01-5P 165314-97-6P 165314-98-7P 165315-02-6P 165315-03-7P 165315-04-8P 165315-05-9P :165315-06-0P 165315-07-1P 165315-08-2P 165315-09-3P 165315-10-6P 165315-11-7P 165315-14-0P 1 165315-15-1P 165315-12-8P 165315-13-9P 165315-16-2P 165315-18-4P 165315-19-5P 165315-20-8P 165315-21-9P 165315-17-3P -165315-24-2P + 165315+25-3P + 165315-26-4P165315-23-1P 165315-22-0P 165315-28-6P (165315-29-7P\*) 165315-30-0P (165315-31-1P 165315-27-5P 165315-32-2P 165315<sup>2</sup>33-3P 165315-34-4P 165315-35-5P 165315-36-6P 三罐料,1997年19月1日中间,1986年 165315-37-7P 165315-38-8P RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of peptide analogs as retroviral protease inhibitors) 建海岸的一块的工厂接入了的工具直接工作工具 ΙT 144114-21-6, Retropepsin

RL: BPR (Biological process); BFOL (Biological study); PROC (Process) (prepn. of peptide analogs as retroviral protease inhibitors) 60-35-5, Acetamide, reactions 7:62-55-5, Thioacetamide 62-56-6, IT Thiourea, reactions 63-91-2, Phenylalanine, reactions 70-23-5, Ethyl bromopyruvate 74-89-5, Methylamine, reactions 75-04-7, Ethanamine, 75-05-8, Acetonitrile, reactions 75-12-7, Formamide, reactions 75-44-5, Carbonic dichloride 78-81-9, Isobutylamine 79-05-0, Propionamide 88-09-5 2-Ethylbutyric acid 105-39-5, Ethyl 107-10-8, 1-Aminopropane, reactions 109-89-7, reactions chloroacetate 

**建构的**与中心。

"最低数算工"的"16"。 4.3 13.54 [10] [16] [16] [17]

```
110-91-8, Morpholine, reactions 122-51-0 123-39-7, N-Methylformam 123-75-1, Pyrrolidine, reactions 503-74-2, 3-Methylbutyric acid 534-07-6, 1,3-Dichloroacetone 539-74-2, Ethyl 3-Dromopropionate 563-83-7, Isobutyramide 638-07-3, Ethyl 4-chloroacetoacetate 867-870-46-2, tert-Butyl carbazate 1118-02-1, Trimethylsilyl isocyanate
                                                  123-39-7, N-Methylformamide
1759-53-1, Cyclopropanecarboxylic acid 2491-20-5, Alanine methyl ester
                  3400-45-1, Cyclopentanecarboxylic acid 3721-95-7,
hvdrochloride
                                5470-11-1 6065-82-3, Ethyl diethoxyacetate
Cyclobutanecarboxylic acid
             6160-65-2, Thiocarbonyl diimidazole 6306-52-1, Valine methyl
6089-04-9
ester hydrochloride / 6372-14-1/2/Phenylalaninol 6921-34-2,
                              6972-05-0, N,N-Dimethylthiourea
                                                                    7204-46-8,
Benzylmagnesium chloride
N, N-Diethylthiourea 13734-41\frac{7}{4}3 16332\frac{1}{4}06-2 16982-21-1, Ethyl
               19967-55-6
                              38585-74-9, 5-Thiazolemethanol
                                                                    65386-28-9
thiooxamate
                              133047-44-6; 154212-61+0 1
165315-47-9 165315-59-3
              131052-44-3
                                                              162739-63-1
65815-64-7
               162740-04-7
                                                               165315-76-4
162739-94-8
                               165315-91-3 165316-16-5
165315-78-6
               165315-80-0
                                                               165316-37-0
RL: RCT (Reactant)
    (prepn. of peptide analogs as retroviral protease inhibitors)
115-08-2P, Methanethioamide 541-46-8P 593-75-9P
                                                             631-58-3P,
Propanethioamide 1114-38-1P 1503-98-6P, Cyclobutanecarboxamide 3217-94-5P, Cyclopentanecarboxamide 6228-73-5P, Cyclopropanecarboxamide
                13515-65-6P, Thioisobutyramide 114294-10-1P,
13242-92-7P
                                15536-75-1P 16536-95-1P 20295-34-5P, 32955-21-8P 32955-22-9P 33142-21-1P
4-Morpholinecarbothioamide
Cyclopropanecarbothicamide
               40398-36-5P, 1-Pyrrolidinecarbothioamide 42202-73-3P,
39624-97-0P
                                 56012-38-5P (59830-60-3P, Z-Phenylalaninal
Cyclopentanecarbothioamide
                98019-60-4P, 5-Isokazolemethanol 98278-52-5P
                                                                       99805-29-5P
79836-78-5P
                                 1111138-83-1P 118994-86-8P,
                 110600#55-0P
104336-01-8P
                                                               126534-22-3P
                             126533 \div 95 \div 7P 126533 \div 96 \div 8P
5-Oxazolecarboxaldehyde
                 127232-41-1P, 5\frac{\pi}{4}0xazolemethanol\frac{1}{4} 133047-45-7P
126534-23-4P
                                 134878-06-1P 134878-07-2P
                                                                    135207-10-2P
                 133333-27-4P
133047-46-8P
                                 143838-10-2P 144141-68-4P 144163-97-3P 144164-10-3P;
                 143747+06-2P
                                                                    144163-44-0P
137649-69-5P
                                                                    144164-11-4P
144163-85-9P
                 144163-96-2P
                                 154212-59-6P 154212-60-9P
                                                                    154248-99-4P
144186-59-4P
                 148682-88-6P
                                                              156732-13-7P
156589-97-8P, Cyclobutanecarbothicamide 156732-12-6P
                 156769-85-6P 162537-10-2P 162739-62-0P
156732-15-9P
                                 162739-70-0P 162739-71-1P
                                                                   :162739-72-2P
162739-68-6P
                 162739-69-7P
                                 162739-99-3P 162740-00-3P
                                                                    162740-01-4P
162739-93-7P
                 162739-98-2P
162740-02-5P
                 162849-92-5P
                                  162849-93-6P 162849-94-7P
                                                                    162849-95-8P
                                                   165315-40-2P
                                                                    165315-41-3P
162849-96-9P
                 162990-03-6P
                                  165315-39-9P
                 165315-43-5P
                                  165315-44-6P
                                                   165315-45-7P
                                                                    165315-46-8P
165315-42-4P
                 165315-49-1P
                                 165315-50-4P 165315-51-5P
                                                                    165315-52-6P
165315-48-0P
                 165315-54-8P
                                 165315-55-9P 165315-56-0P
                                                                    165315-57-1P
165315-53-7P
                                 165315-61-7P 165315-62-8P
                                                                    165315-63-9P
                 165315-60-6P
165315-58-2P
                                  165315-66-2P 165315-67-3P
                                                                    165315-68-4P
165315-64-0P
                 165315-65-1P
                                  165315-71-9P 165315-72-0P
                 165315-70-8P
                                                                    165315-73-1P
165315-69-5P
                                  165315-77-5P 165315+79-7P
165315-74-2P
                 165315-75-3P
                                                                    165315-81-1P
                                165315-84-4P 1 165315-85-5P
                 165315-83-3P
165315-82-2P
                                                                    165315-86-6P
                                                                   165315-92-4P
165315-87-7P
                 165315-88-8P
                                  165315-89-9P - 165315-90-2P
                                  165315-95-7P 165315+96-8P
                                                                    165315-97-9P
165315-93-5P
                165315-94-6P
                                  165316-00-7P 165316+01-8P
                                                                   [165316-02-9P
                 165315-99-1P
165315-98-0P
                                 165316-05-2P
                                                   165316-06-3P
                                                                    165316-07-4P
                 165316-04-1P
165316-03-0P
                                                 165316+11-0P
                                                                    165316-12-1P
                 165316-09-6P
                                  165316-10-9P
165316-08-5P
                                 165316-15-4P 165316+17-6P
                                                                    165316-18-7P
165316-13-2P
                 165316-14-3P
                                                   165316-22-3P
                                                                    165316-23-4P
                                 165316-21-2P
165316-19-8P
                 165316-20-1P
                                 165316-26-7P 165316-27-8P
                                                                    165316-28-9P
165316-24-5P
                 165316-25-6P
                                 165316-31-4P 165316-32-5P
                                                                    165316-33-6P
165316-29-0P
                 165316-30-3P
                                  165316-36-9P* 165316-38-1P
                                                                    165316-39-2P
165316-34-7P
                 165316-35-8P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
    (prepn. of peptide analogs as retroviral protease inhibitors)
```

Aug Traffi

1984 14 B M

1.15本股份以前

744 15

Gundalte di 1921

14 to 1

IT

ΙT 65815-64-7

RL: RCT (Reactant)

(prepn. of peptide analogs as retroviral protease inhibitors)

RN65815-64-7 HCAPLUS

L-Alanine, N-[(4-nitrophenoxy) carbonyl]-, methyl ester (9CI)
NAME) CN (CA INDEX

Absolute stereochemistry.

IT

162537-10-2P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)

RN

(prepn. of peptide analogs as retroviral professe inhibitors)
162537-10-2 HCAPLUS
L-Valine, N-[(4-nitrophenoxy)carbonyl], methylester (9CI) (CA CN 1. 据题中分别的 NAME)

医新生菌素

Absolute stereochemistry.

ANSWER 16 OF 24 HCAPLUS COPYRIGHT 2001 ACS L27

11

AN 1995:522611 HCAPLUS

DN 122:291525

Retroviral protease inhibiting compounds ΤI

Norbeck, Daniel W.; Sham, Hing Leung; Kempf, Dale J.;
Abbott Laboratories, USA
PCT Int. Appl., 185 pp.
CODEN: PIXXD2
Patent
English IN Zhao, Chen

PA

SO

DT

LΑ English

IC

DAMENIM NO

ICM C07D263-34 | A61K031-42; A61K031-425,

CC 34-2 (Amino Acids, Peptides, and Aller 1997) Proteins)

Section cross-reference(s): 1

FAN.CNT 2

	PATENT NO.	KIND DATE (APPLICATION NO. DATE	
ΡI	WO 9419332 W: CA, JP	A1 19940901 WO 1994-US1457 19940208	
	•	CH, DE, DK, ES, FR; GB, GR, IE, IT, LU, MC, NL, PT, SE	
	US 5461067	A 1 1 19951024 1 Us 1994-185666 19940201	
	EP 683772	A1 19951129 EP 1994-908018 19940208	
	R: AT, BE,	CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE	

T2 JP 08507061 19960730 JP 1994-519025 19930225 PRAI US 1993-23226 1 19940201 US 1994-185666 WO 1994-US1457 19940208 MARPAT 122:291525 os GΙ 

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

A PROPERTY OF

Retroviral protease-inhibiting compds. are disclosed, specifically I [R1, AB R2 = H, (un) substituted alkyl, aryl, alkenyl, heterocyclyl, substituted carbonyl; Y = NHCHR4CO, NHNR4CO, Q1, Q2; Y' = COCHR3NH, CONR3NH, Q3, Q4; a, b = 0-3; c, d = 1-2; R3, R4 = H, (un) substituted alkyl, aryl, alkenyl, heterocyclyl, substituted carbonyl; R3', R4' = H, alkyl; m, n = 0-1; R5, R6 = C(T)GR7; T = O, S; G = CH2, O, S, NR8; R7 = (un)substituted alkyl or cycloalkyl, aryl, protecting group; <math>R8 = H, alkyl, cycloalkyl] and their salts, esters, and prodrugs. For example, 5-(hydroxymethyl)thiazole (prepd. in 4 steps) reacted with 4-nitrophenyl chloroformate to give 78% of the corresponding carbonate. This reacted with 2-(tert-butoxycarbonylamino)-4S-hydroxy-5S-amino-1,6-diphenyl-2-azahexane (prepn. given), followed by deprotection and coupling with a corresponding valine deriv., to give title compd. II. At 0.5 nM in vitro, II gave 77% inhibition of HIV-1 protease. II also gave 50% inhibition of cytopathy of MT4 cells by HIV-13B at 0.10-0.11 .mu.M, with a cellular LC50 of 17.mu.M. Examples include 69 syntheses (some prophetic), and similar biol. data for other selected I.

valinyl aminohydroxyazahexane prepn retroviral protease inhibitor; HIV ST protease inhibitor aminohydroxyazahexane prepniliti

Virucides and Virustats ΙT

> (prepn. of valinylaminohydroxyazahexane derivs. and analogs as retroviral protease inhibitors)

Peptides, preparation IT

> RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES 131,51

一个连续的第三词 的复数加快

(prepn. of valinylaminohydroxyazahexane derivs and analogs as retroviral protease inhibitors)

ΙT Virus, animal

(human immunodeficiency, prepn. of valinylaminohydroxyazahexane derivs. and analogs as retroviral protease inhibitors) 一种 自身种种

144114-21-6, Retropepsin IT

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (HIV; prepn. of valinylaminohydroxyazahexane derivs. and analogs as retroviral protease inhibitors)

60222-90-4P, 5-Hydroxypentanal oxime 128018-44-0P ΙT

111

RL: BYP (Byproduct); PREP (Preparation)

(byproduct; prepn. of valinylaminohydroxyazahexane derivs. and analogs as retroviral protease inhibitors)

115-08-2P, Thioformamide 934-53-2P, 2-Chloro-2-phenylpropane IT 13242-92-7P, 4-(Chloromethyl)-2-(dimethylamino)thiazole 13515-65-6P, 2-Methylpropanethioamide 14294-10-1P, 4-(Aminothiocarbonyl) morpholine 15536-75-1P, 2-Methoxythioacetamide 16689-35-3P 24469-50-9P 30293-86-8P, .alpha. ~Isocyanatovaline methyl ester 32939-32-5P 32955-22-9P, Ethyl thiazole-5-carboxylate 33142-21-1P, Ethyl 2-chloro-2-formylacetate 38585-74-99, 5-(Hydroxymethyl)thiazole 39624-97-0P 40398-36-5P, 1-[Amino(thiocarbonyl)]pyrrolidine हिस्सामा स्टार्च र माल व्हे के हैं है। या व

man 豆腐物 基位。

Marie Caracillo Del Victoria de la compania de la compania del Caracillo Del Caracillo

```
41337-78-4P, 2-Carboethoxy-6-ethylpyridine 53370-84-6P 57699-48-6P
                   59830-60-3P, N-Benzyloxycarbonyl-L-phenylalaninal
     57699-55-5P
     65386-28-9P, 4-(Chloromethyl)-2-isopropylthiazole hydrochloride
     98019-60-4P, 5-(Hydroxymethyl) isoxazole 99805-29-5P
                                                              100868-72-2P,
     2-[(N-Methylamino)methyl]pyridine dihydrochloride 126533-95-7P, Ethyl
     2-(4-morpholinyl)thiazole-4-carboxylate 126533-96-8P,
     2-(4-Morpholinyl)-4-(hydroxymethyl)thiazole .134807-06-0P
                                                                     134807-20-8P
                                  134807-30-0P 137515-66+3P
144186-53-8P 144186-59-4P
                   134807-29-7P
                                                                    144163-45-1P
     134807-28-6P
                                                                    149267-56-1P
     144163-71-3P
                    144163-97-3P
                                    149267-73-2P 150767-00-3P
                                                                   150767-02-5P
     149267-64-1P 149267-65-2P
                                   150767-08-1P
     150767-04-7P
                    150767-05-8P
                                                  154212-60-9P
                                                                    154212-61-0P
     154248-99-4P 155884-24-5P, Methyl 2-isopropyl-4-oxazolecarboxylate
    161852-61-5P 162537-10-2P 162739-61-9P 162739-62-0P 162739-63-1P 162739-64-2P 162739-65-3P 162739-66-4P 162739-71-1P 162739-76-6P 162739-77-7P 162739-78-8P
                                                                    162739-67-5P
     162739-71-1P
                                                                    162739-79-9P
     162739-80-2P
                    162739-81-3P
                                    162739-82-4P
                                                    162739-83-5P
                                                                    162739-84-6P
                                    162739-93-7P 162739-94-8P,
     162739-85-7P
                    162739-86-8P
                                                                      162739-96-0P
     2-Isopropyl-4-[(N-methylamino)methyl]oxazole 162739-95-9P
     162739-97-1P, 2-[(N-Methylamino)methyl]-6-ethylpyridine | 162739-98-2P,
     2-(N,N-Dimethylamino) 4-(hydroxymethyl)thiazole 162739-99-3P, Ethyl
     2-(1-pyrrolidinyl)thiazole-4-carboxylate 162740-00-3P,
     2-(1-Pyrrolidinyl)-4-(hydroxymethyl) thiazole 162740-01-4P,
     4-(Chloromethyl)-2-(methoxymethyl)thiazole@hydrochloride 162740-02-5P,
     2-(Methoxymethyl)-4-["N-methylämino)methyl]thiazole 162740-03-6P,
     4-Hydroxymethyl-2-isopropyloxazole 163658-33-1P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (intermediate; prepn. of valinylaminohydroxyazahexane derivs. and
        analogs as retroviral protease inhibitors)
                                    162739-68-6P 162739-69+7P
162739-74-4P 162739-75+5P
                                                    162739-69-7P
                                                                    162739-70-0P
ΙT
                    144186-54-9P
     144163~72-4P
                    162739-73-3P
     162739-72-2P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (intermediate; prepn. of valinylaminohydroxyazahexane derivs. and
        162739+24-4P 1162739-28-8P.
                                                                    162739-31-3P
                    162739<del>-</del>20-0P
IT
     150767-06-9P
     RL: ADV (Adverse effect): including toxicity); BAC (Biological activity or
     effector, except adverse); RCT \(Reactant); SPN (Synthetic preparation);
     THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
                          27 11
     (Uses)
        (prepn. of valinylaminohydroxyazahexane derivs. and analogs as retroviral protease inhibitors)
                                    162739-26-6P 162739-27-7P
IT
     162739-22-2P
                    162739-25-5P
                                                                    162739-29-9P
     162739-30-2P 162739-32-4P 3 BAC (Biological activity or
     effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic
     use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of valinylaminohydroxyazahexane derivs. and analogs as
        (prepn. of Valiny and Honya. 1973) retroviral protease inhibitors) 767-07-0P 150767 09-2P 162739 33-5P 162739 35-7P
IT
     150767-07-0P
                                                                    162739-38-0P
     162739-43-7P
                     162739-46-0P
     RL: BAC (Biological activity or effector, except adverse); RCT (Reactant);
     SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of valinylaminohydroxyazahexane derivs and analogs as
        retroviral protease inhibitors)
                    162739+21-1P 162739+23-3P 162739-34-6P 162739-36-8P 162739-3P 162739-40-4P 162739-41-5P 162739-42-6P
IT
     150767-10-5P
     162739-37-9P
                                    162739-47-1P 162739-48-2P 162739-49-3P
                    162739-45-9P
     162739-44-8P
                                    162739-52-8P 112739-53-9P
                                                                  162739-54-0P
                    162739-51-7P
     162739-50-6P
                                   162739-59-5P
                    162739-56-2P
     162739-55-1P
     162739-60-8P
```

理信仰 计对应

RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOI (Biological study); PREP (Preparation); USES (Uses)
(prepn. of valinylaminohydroxyazahexane derivs; and analogs as retroviral protease inhibitors) 67-64-1, Acetone, reactions 70-23-5 Ethyl bromopyruvate 74-89-5, ΙT Methylamine, reactions 75-12-7, Formamide, reactions 75-44-5, Carbonic 98-01-1, 2-Furaldehyde, reactions 98-83-9; .alpha.-Methylstyrene; reactions 100-52-7, Benzaldehyde, reactions 100-55-0, Pyridine-3-methanol 100-83-4, 3-Hydroxybenzaldehyde 105-39-5, Ethyl chloroacetate 109-94-4, Ethyl formate 110-91-8, Morpholine, reactions 122-51-0, Triethyl orthoformate 123-08-0, 4-Hydroxybenzaldehyde 123-11-5, p-Anisaldehyde, reactions 123-7 Pyrrolidine, reactions 459-57-4, 4-Fluorobenzaldehyde 498-60-2, 534-07-6, 1,3-Dichloroacetone 563-83-7, Isobutyramide 3-Furaldehyde 586-98-1, Pyridine-2-methanol 591-31-1, m-Anisaldehyde 688-3-Hydroxy-5-hexene 870-46-2, tert-Butyl carbazate 872-85-5, Pyridine-4-carboxaldehyde 925-90-6, Ethylmagnesium bromide 688-99-3, 930-45-0, 1121-60-4, Pyridine-2-carboxaldehyde (S,S)-2-Aminocyclopentanol 1779-49-3, Triphenylmethylphosphonium bromide 2043-61-0, Cyclohexanecarboxaldehyde 3731-51-9, 2-(Aminomethyl)pyridine 5470-11-1, Hydroxylamine hydrochloride 6089-04-9, 3,4,5,6-Tetrahydro-2-(2-propynyloxy)-2H-pyran 6160-65-2, Thiocarbonyl diimidazole 6306-52-1, L-Valine methyl ester hydrochloride 6372-14-1, N-Benzyloxycarbonyl-L-phenylalaninol 6972-05-0, N,N-Dimethylthiourea 7664-41-7, Ammonia, reactions 37693-46-1, 4-Nitrophenyl chloroformate 14337-43-0, Ethyl chlorooximidoacetate 16332+06-2, 2-Methoxyacetamide 74111, 21 0, (S, 5) -2 - Aminocyclohexanol 57699-57-7 69353-16-8 82625-45-4, 4-(2-Morpholinoethoxy) benzaldehyde 130782-46-6 135941-95-6 162739-8880 162739-89-1 (162739-90-4 135941-97-8 162739#87+9 162740-04-7, 4-(Chloromethyl)-2-162739-92-6 162739-91-5 (dimethylamino) thiazole dihydrochloride 162740 705 78, N Isobutyrylserine 34. 数16年的 11 11/1 methyl ester THE WAR THE WAR RL: RCT (Reactant) (reactant; prepn. of valinylaminohydroxyazahexane derivs; and analogs

as retroviral protease inhibitors)

## ΙT 162537-10-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (intermediate; prepn. of valinylaminohydroxyazahexane derivs. and

"提展情 

RN

analogs as retroviral protease inhibitors)

162537-10-2 HCAPLUS

L-Valine, N-[(4-nitrophenoxy)carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

ute stereochemistry. CN

Absolute stereochemistry.

ANSWER 17 OF 24 HCAPLUS

· High

AN 1991:186008 HCAPLUS

DN

Synthesis and spectroscopic properties of azaglutamine amino acid and ΤI FOREILIES OF ALL STATES peptide derivatives

```
Gray, C. J.; Quibell, M.; Jiang, K. L.; Baggett, N.
AU
     Sch. Chem., Univ. Birmingham, Birmingham, B15 2TT,
CS
     Synthesis (1991), (2), 141-6
SO
     CODEN: SYNTBF; ISSN: 0039-7881
     Journal
DT
                                      各的多道市基份工作。(1)
     English
LΑ
CC
     34-3 (Amino Acids, Peptides, and
     Proteins)
     CASREACT 114:186008
os
     Azaglutamines H2NCOCH2CH2N (NHBoc) CO2R (Boc = Me3CO2C; R = Et, CH2Ph) were
AΒ
     prepd. by treating H2NCOCH2NHNHBOc (I) with ClCO2R. Dipeptides
     H2NCOCH2CH2N (NHBoc) CO-X-OCH2Ph (X = Gly, Phe) were prepd. by treating I with 2,4-(O2N) C6H3O2C-X-OCH2Ph I was treated with OCNCH2CO2Et to give H2NCOCH2CH2N (NHBoc) CONHCH2CO2Et
     azaglutamine peptide; glutamine aza
ST
     Peptides, preparation
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
         (azaglutamine-contg., prepn. and NMR of) 🕟
                                                      41863-52-9
     870-46-2, tert-Butyl carbazate 330189-48-1
IT
         RCT (Reactant)
(addn. reaction of, with acrylamide)
     RL: RCT (Reactant)
     79-06-1, 2-Propenamide, reactions
RL: RCT (Reactant)
IT
         (addn. reaction of, with tert-Bu carbazate)
                                     133382-92-0P 133382-95-3P
                     133382-91-9P
                                                                    133382-96-4P
IT
     133382-90-8P
                                    133383-04-7P 133383-05-8P
                                                                     133383-06-9P
     133382-99-7P
                     133383-00-3P
     133383-07-0DP, derivs:
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. and NMR of)
IT
     133382-98-6P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. and reaction of, with Et isocyanatoacetate)
     1738-76-7P, Glycine benzyl ester tosylate 1738-78-9P, L-Phenylalanine
IT
                                        Sharangar Shire Parisher and
     benzyl ester tosylate
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. and reaction of, with bis (dinitrophenyl) carbonate)
     133382-89-5P
ΙT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. and reaction of, with chloroformates or Et isocyanoacetafte)
                                       是重新的對於小學也不同數學工作。
                        -1
     133382-97-5P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. and reaction of, with dinitrophenoxycarbonyl amino acid esters)
ΙT
     133382-93-1P 133382-94-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
         (prepn. and reaction of, with hydrazide)
IT
     133383-19-4P
     RL: SPN (Synthetic preparation) PREP (Preparation)
         (prepn. of)
IT
                     133383-02-5P 133383-03-6P
     133383-01-4P
     RL: SPN (Synthetic preparation); PREP (Preparation)
     (prepn., sapon. and NMR of) 501-53-1, Benzyl chloroformate
ΙT
     7497-12-3, Bis(2,4-dinitrophenyl) carbonate
                                    相解: [[1] [1] [1] [1] [1]
     RL: RCT (Reactant)
         (reaction of, with (tert-butoxycarbonylhydraino)propanamide)
     2949-22-6, Ethyl isocyanato acetate RL: RCT (Reactant)
IT
         RCT (Reactant)
(reaction of, with hydrazides)
IT
     133382-93-1P 133382-94-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
```

翻 回语的特性 . 种种的生物的

100

(prepn. and reaction of, with hydrazide)

RN 133382-93-1 HCAPLUS

CN L-Phenylalanine, N-[(2,4-dinitrophenoxy)carbonyl]-, phenylmethyl ester

(9CI) (CA INDEX NAME)

Absolute stereochemistry:

RN 133382-94-2 HCAPLUS CN Glycine, N-[(2,4-dinitrophenoxy)carbonyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

$$O_{2}$$
  $O_{1}$   $O_{2}$   $O_{2}$   $O_{2}$   $O_{3}$   $O_{4}$   $O_{2}$   $O_{4}$   $O_{5}$   $O_{7}$   $O_{7$ 

ANSWER 18 OF 24 HCAPLUS COPYRIGHT 2001 ACS L27 1991:62698 HCAPLUS ΑN : 1 th : 1 to a DN New macrocyclic pseudopeptides containing urethane backbone linkages ΤI Wu, Youling; Kohn, Joachim ΑU Dep. Chem., Rutgers, State Univ., New Brunswick, NJ, CS J. Am. Chem. Soc. (1991), 113(2), 687-8 SO CODEN: JACSAT; ISSN: 0002-7863 DT Journal English LΑ 34-3 (Amino Acids, Peptides, and CC Proteins) OS CASREACT 114:62698 GΙ

- \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT
- The use of urethane bonds as a new type of amide bone mimetic in the design of pseudopeptides was investigated. Urethane backbone linkages were derived from the side chain hydroxyl group of tyrosine by reacting Boc-Tyr-OHex (Boc = Me3CO2C, Hex = hexyl) with p-nitrophenyl chloroformate, followed by reaction with alanine benzyl ester to give urethane-bonded pseudodipeptide I. The resulting Tyr-Ala pseudodipeptide could be cyclized to 24- and 36-membered macrocyclic pseudopeptides II and III, resp., with cyclization yields of about 80%. The unusual ease with which the Tyr-Ala pseudodipeptide underwent cyclizations is probably a consequence of the cis conformation of the urethane bond. In a phase

```
transfer expt., the ion binding properties of the 24-membered macrocycle
     II were studied. II was an effective and selective phase transfer agent
     for Li+ ions, solubilizing up to 0.6 molar equivalents of Li+ ions in
     chloroform. The uptake of Na+ and K+ ions was only 0.025 and 0.007 molar equivalents, resp. Since the structure of II can be readily modified, the
    synthetic approach gives rise to a family of new macrocyclic pseudopeptides.
macrocyclic pseudopeptide urethane backbone linkage
Alkali metals, reactions
RL: RCT (Reactant)
st
IT
     RL: RCT (Reactant)
        (binding of, with macrocyclic pseudopeptide contg. urethane backbone
        linkages)
IT
     Peptides, preparation
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (cyclopseudo-, prepn. of, with urethane backbone linkages)
                3324-58-1 18390-55-1 Process)
IT
     573-83-1
     RL: PROC (Process)
        (binding of, with macrocyclic pseudopeptide contg. urethane backbone
        linkages)
                                    "工具的制造技术"。
IT
     17831-01-5
     RL: RCT (Reactant)
        (coupling of, with tyrosine active carbonate deriv.)
IT
     131152-83-5P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and binding of, with alkali metals)
ΙT
     131152-91-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and coupling of, with alanine benzyl ester)
IT
     131152-89-1P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and cyclization of)
                                       ΙT
     131152-86-8P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and esterification of, with nitrophenol)
152-87-9P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation);
        (prepn. and partial deblocking of)
     131152-90-4P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and reaction of, with nitrophenyl chloroformate)
     131152-84-6P
ΙT
     RL: SPN (Synthetic preparation); PREP (Preparation)
                                    (prepn. of)
     131152-85-7P
IT
     RL: SPN (Synthetic preparation): PREP (Preparation)
        (prepn., catalytic transfer hydrogenolysis, and metal-binding
        properties of)
     7693-46-1, p-Nitrophenyl chloroformated in the
ΙT
     RL: RCT (Reactant)
        (reaction of, with tyrosine deriv.)
                                      IT
     94326-61-1
     RL: RCT (Reactant)
        (tert-butoxycarbonylation of)
IT
     131152-89-1P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and cyclization of)
                                    李基斯 中的时间 "我们的"我们的我们的"。
RN
     131152-89-1 HCAPLUS
     L-Tyrosine, hexyl ester, ester with N-carboxy-L-alanine 1-(4-nitrophenyl)
CN
     ester, mono(trifluoroacetate) (9CI) (CA INDEX NAME)
```

CM 1

CRN 131152-88-0 CMF C25 H31 N3 O8 CDES \*

CM 2

CRN 76-05-1 CMF C2 H F3 O2

IT 131152-87-9P

RL: SPN (Synthetic preparation); PREP (Preparation (prepn. and partial deblocking of)

RN 131152-87-9 HCAPLUS

CN L-Tyrosine, N-[(1,1-dimethylethoxy) carbonyl] +, hexyl ester, ester with N-carboxy-L-alanine 1 (4-nitrophenyl) ester (9CI) (CA INDEX NAME)

L27 ANSWER 19 OF 24 HCAPLUS COPYRIGHT 2001 ACS

AN 1990:99261 HCAPLUS

DN 112:99261

TI Preparation of N-(phosphonocyclohexylhydroxypropyl) derivatives of amino acids and dipeptides as renin inhibitors

IN Patel, Dinesh V.

PA Squibb, E. R., and Sons, Inc., USA

SO Eur. Pat. Appl., 121 pp.

CODEN: EPXXDW

DT Patent

LA English

IC ICM C07K005-06

A61K037-64; C07F009-40; C07F009=65; C07F009+32; C07F009-44; 4:513 A61K031-66 34-3 (Amino Acids, Peptides, and Section cross-reference(s): 1 FAN.CNT 1 KIND, APPLICATION NO. DATE PATENT NO. EP 1989-103489 19890906 EP 331105 A2 · 19890228 19900905 **A3** EP 331105 R: AT, BE, CH, DE, ES, FR, GB, IT, LI, LU, NL, SE **A1** : 19890223 19890908 WO 1989-US777 WO 8907940 W: DK, HU, JP HU 1989-2302 A2 19900828<sup>2</sup> 19890223 HU 52785 JP 02503440 T2 19901018 JP 1989-503323 19890223 ZA 8901594 Α 19891129 ZA 1989-1594 19890301 AU 1989-30999 AU 8930999 A1 19890907 19890302 Α DK 1989-5469 DK 8905469 19891227 19891102 Α 19930608 US 1990-509398 19900412 US 5217958 19880303 PRAI US 1988-163593 WO 1989-US777 19890223 US 1989-317257; 19890228 MARPAT 112:99261 alkyl) carbamoyl or -sulfamoyl, substituted alkanoyl, alkoxycarbonyl, alkylthio, alkanoylthio, etc.; when Y = 0, X = N-(substituted)when Y = NH, X = N-(substituted alkyl) carbamoyl, substitutedalkoxycarbonyl, alkanoyl, alkyl, alkylthio, etc.; R1, R2 = H,

OS Amino acid and dipeptide derivs: X-Y-(CHR5CONH)rCHR4CONHCHR3CH(OH)P(:M)(Z1 AB R2) ZR1 [I; M = O, S; Y = CH2, NH, O; when Y = CH2, X = N-(substituted alkyl)carbamoyl, substituted alkanoyl, alkoxycarbonyl, phosphono, etc.; (cyclo) alkyl, arylalkyl, (hetero) aryl; Z, Z1 = bond, (un) substituted CH2, NH, cyclic amino, heterocyclyl; R3, R5 = H, lower (halo)alkyl, arylalkyl, heterocyclylalkyl, cycloalkyl, (CH2) nOH, (CH2) nNH2, (CH2) nSH, (CH2) nNHC(:NH) NH2, (CH2) nCONH2, N-substituted 5-imidazolylalkyl, etc.; R4 = any group defined for R3 and R5, N-substituted 2-imidazolyl, 4- or 2-thiazolylalkyl, 3-pyrazolylalkyl, 4-for 2-oxazolylalkyl; n = 1-5], useful as cardiovascular agents in the treatment of hypertension, congestive heart failure, renin-dependent hyperaldosteronism, myocardial infarction, etc., and as a diagnostic agent in detg. renin related disorders (no data), were prepd. I are also inhibitors of retroviral protease and thus are useful as virucides against human T-cell leukemia virus HTLV-1 and HTLV-III (no data). Thus, hydrogenation of (S)-BOC-Phe-OH (BOC = Me3CO2C) over PtO2 and amidation of the resulting (S)-BOC-NHCHQCO2H (Q = cyclohexylmethyl) with MeNHOMe.HCl in THF contg. carbodimidazole gave (S)-BOC-NHCHQCONMeOMe. Redn. of the latter with LiAlH4 in THF/Et20 at 0.degree to (S)-BOC-NHCHQCHO followed by addn. reaction with (MeO) 2PH in DMF in the presence of KF gave a 12.7:1.0 diastereomeric mixt. of (1S)-BOC-NHCHQCH(OH)P(OMe)2 (Q unchanged). N-Deprotection of the latter with 1.2 N HCl in EtOAc and condensation with BOC-Leu-OH.H2O in DMF in the presence of hydroxybenzotriazole, Et3N, and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide-HCl gave a major diastereomer (1S)-BOC-Leu-NHCHQCH(OH)P(OMe)2. Approx. 25 I were prepd. ST amino acid phosphonocyclohexylhydroxypropyl renin inhibitor; peptide phosphonocyclohexylhydroxypropyl prepni renin inhibitor; cardiovascular agent phosphonocyclohexylhydroxypropyl dipeptide amide; antihypertensive phosphonocyclohexylhydroxypropyl dipeptide amide: 11 11 IT Aldosteronism

(treatment of, N-(phosphonohydroxyalkyl)amino acids and -dipeptides 4月7日 新广东部 1.7 1 1 - 1 **操的**通信。 加克基基

THE PROPERTY OF STREET

**明顯於自然生化。1945年前以上** 

Antihypertensives IT

Butterly

126

CC

ΡI

```
Cardiovascular agents
         Virucides and Virustats
               (N-(phosphonohydroxyalkyl) amino acid and dipeptide amides)
ΙT
         Amides
         RL: SPN (Synthetic preparation); PREP (Preparation)
               (amino, N-(phosphonohydroxyalkyl), preph. of, as renin inhibitors and
               cardiovascular agents)
ΙT
         Peptides, compounds
         RL: SPN (Synthetic preparation); PREP (Preparation)
               (di-, amides, N-(phosphonohydroxyalkyl), prepn. of, as renin inhibitors
               and cardiovascular agents)
IT
               (failure, treatment of, N-(phosphonohydroxyalkyl)amino acids and
                                                                   建筑市的
               -dipeptides for)
         Heart, disease or disorder
ΙT
                (infarction, treatment of, N-(phosphonohydroxyalkyl)amino acids and
               -dipeptides for)
                                                                   [4] · [1] · [1] · [1] · [1] · [1] · [1]
IT
         3587-60-8
                                                                  AP Shipst Mark the
         RL: RCT (Reactant)
         RL: RCT (Reactant)
(N-alkylation by, of histidine Me ester deriv.)
7693-46-1, p-Nitrophenyl chloroformate
ΙT
         RL: RCT (Reactant)
               (acylation by, of Me phenylalaninate)
         (acylation by, of me phenylers, 501-53-1, Benzyl chloroformate
IT
         RL: RCT (Reactant)
         (acylation by, of aminocaproic acid)

108-23-6, Isopropyl chloroformate 1070-83-3, tert-Butylacetic acid

RL: RCT (Reactant)
IT
                (acylation by, of histidinamide deriv.)
         24424-99-5, Di-tert-Butyl dicarbonate graduate of the state of the sta
IT
               RCT (Reactant)
(acylation by, of histidine Me ester)
         RL: RCT (Reactant)
IT
         645-45-4, Hydrocinnamoyl chloride 👍
               RCT (Reactant)
(acylation by, of leucine)
         RL: RCT (Reactant)
         3400-45-1, Cyclopentanecarboxylic acid
IT
                                                 The Manual in the Lord is a
         RL: RCT (Reactant)
               (acylation by, of phenylalanylleucinamide)
IT
         60-32-2
                                               RL: RCT (Reactant)
                (acylation of, by benzyl chloroformate)
         7524-50-7, L-Phenylalanine methyl ester hydrochloride
IT
         RL: RCT (Reactant)
               (acylation of, by cyclopentanecarboxylic acid)
         61-90-5, L-Leucine, reactions
IT
         RL: RCT (Reactant)
               (acylation of, by hydrocinnamoyl chloride)
         762-04-9 868-85-9, Dimethyl phosphite 1990 1991
ΙT
         RL: RCT (Reactant)
                (addn. reaction of, with aminocyclohexylpropanal deriv.)
         75-26-3, 2-Bromopropane 78-77-3, 1-Bromo-2-methylpropane RL: RCT (Reactant)
IT
                (alkylation by, of [(hydroxymethoxyphosphinyl)ethyl]oxazolidinecarboxyl
               ate)
         25024-53-7
IT
         RL: RCT (Reactant)
                (amidation of, with (aminophenyl) phosphonate deriv.)
                                                                  IT
         2018-66-8
         RL: RCT (Reactant)
                (amidation of, with (aminopropyl)phosphonate deriv.)
```

Angele and Angele and

MAN THE STATE OF THE

```
110-91-8, Morpholine, reactions
IT
     RL: RCT (Reactant)
         (amidation of, with Me [(nitrophenoxy)carbonyl]phenylalaninate)
     6638-79-5, O,N-Dimethylhydroxylamine hydrochloride
IT
     RL: RCT (Reactant)
         (amidation of, with aminocyclohexylpropanoic acid deriv.)
     501-52-0, Hydrocinnamic acid RL: RCT (Reactant)
IT
     RL: RCT (Reactant)
         (amidation of, with leucinamide deriv.)
     74-89-5, Methylamine, reactions
RL: RCT (Reactant)
(amination by, of (hydroxymethoxyphosphinyl)oxazolidinecarboxylate)
IT
                                       IT
     13139-15-6
     RL: RCT (Reactant)
         (condensation of, with di-Me (aminocyclohexylhydroxypropyl) phosphonate)
     71-00-1, L-Histidine, reactions
ΙT
     RL: RCT (Reactant)
(conversion of, to Me ester)
77-76-9, 2,2-Dimethoxypropane
ΙT
     RL: RCT (Reactant)
         (cyclocondensation of, with [(cyclohexylmethyl)hydroxyethyl]carbamate
     deriv.)
1498-40-4, Ethyl dichlorophosphine
RL: RCT (Reactant)

/--terification of, with methanol)
IT
ΙT
     13734-34-4
RL: RCT (Reactant)
(hydrogenation of)
     9015-94-5, Renin, uses and miscellaneous RL: USES (Uses)
IT.
         (inhibitors, N-(phosphonocyclonexylhydroxypropyl)amino acids and
         dipeptides)
     75-16-1, Methylmagnesium bromide
RL: RCT (Reactant)
IT
         (methylation by, of [(hydroxymethoxyphosphinyl)ethyl]oxazolidinecarboxy
                        late)
     2666-93-5
IT
         RCT (Reactant)
(peptide coupling of, in prepn. of renin inhibitor)
7-00-8P 2752-56-9P 15027-08-4P 20898-43-5P 22888-60-4P,
     RL: RCT (Reactant)
ΙT
     1947-00-8P
     L-Histidine methyl ester hydrochloride, 27852-48-8P 33014-68-5P
     37736-82-6P 38155-45-2P 54601-21-7P 64152-76-7P
                     75691-91-7P 83468-82-0P 83468-83-1P 98105-42-1P
     64155-03-9P
                                                          115766-13-7P 120915-52-8P
                                        114473-20-0P
     111629-40-4P
                      114457-62-4P
                                       125399-33-9P 125399-34-0P 125399-35-1P

    120915-53-9P
    121533-66-2P
    125399-33-9P
    125399-34-0P

    125399-36-2P
    125399+37-3P
    125399+38-4P
    125399-39-5P

    125399-41-9P
    125399-42-0P
    125399-43-1P
    125399-44-2P

    125399-46-4P
    125399-47-5P
    125399-48-6P
    125399-49-7P

                      121533-66-2P 1125399-33-9F 125399-39-5P 125399-40-8P 125399-42-0P 125399-43-1P 125399-44-2P 125399-45-3P 125399-47-5P 125399-48-6P 125399-49-7P 125399-50-0P
                                                                           125399-50-0P
                                        125399-53-3P 125399-54-4P
     125399-51-1P 125399-52-2P
                                                                           125399-55-5P
     125399-56-6P 125399-57-7P 125399-58-8P 125399-59-9P 125435-73-6P 125435-74-7P 125435-75-8P 126431-13-8P
                                                                           125399-60-2P
                                                        126431-13-8P
                                                                           126431-14-9P
     RL: SPN (Synthetic preparation); PREP (Preparation)
         (prepn. of, as intermediate for renin inhibitor and cardiovascular
                       125399-10-2P 125399-12-4P: 125399-13-5P: 125399-14-6P
         agent)
IT
     125399-09-9P
                       125399-16-8P 4125399-17-9P 1.125399-19-1P 125399-20-4P
      125399-15-7P
     125399-21-5P 125399-22-6P 125399-23-7P 125399-24-8P 125399-25-9P
     125399-26-0P
                       125399-27-1P 125399-28-2P 125399-29-3P
                                                                           125399-30-6P
     125399-31-7P 125399-32-8P 125435-72-5P 125472-49-3P
     RL: SPN (Synthetic preparation); PREP (Preparation)
```

A STATE OF THE STA

```
(prepn. of, as renin inhibitor and cardiovascular agent)
```

75-77-4, Trimethylsilyl chloride, reactions RL: RCT (Reactant) IT

RL: RCT (Reactant)

(silylation by, of [(cyclohexylmethyl)(ethylmethoxyphosphinyl)hydroxy amate) ethyl]carbamate)

25.325 43 

IT 54601-21-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of, as intermediate for renin inhibitor and cardiovascular 3464

54601-21-7 HCAPLUS RN

CN L-Phenylalanine, N-[(4-nitrophenoxy)carbonyl]-, methyl ester (9CI) INDEX NAME)

Absolute stereochemistry.

ANSWER 20 OF 24 HCAPLUS COPYRIGHT 2001 ACS 1985:7099 HCAPLUS 102:7099 L27

AN

DN 102:7099

8-Quinolinyl carbamates and their use as urinary tract antimicrobials ΤI

Africally pr

Paxton, Larry D.; Madison, Rita A.; Dunbar, Joseph E.
Dow Chemical Co., USA
U.S., 6 pp.
CODEN: USXXAM
Patent IN

PA

SO

DTPatent

LΑ English

A61K031-47; C07D215-34 ·IC

NCL 424258000

34-3 (Amino Acids, Peptides, and and a restard CC

Proteins)
Section cross-reference(s): 10, 27

FAN.CNT 1

APPLICATION NO. KIND DATE PATENT NO. 19840918 US 1982-408292 US 4472404 19820816 PΙ GI

```
Title compds. I (R1 = H, alkyl; R2 = H, NO2, halo; R3 = H, halo; R4 =
AΒ
     alkyl) were prepd. as title agents. Thus, quinoline II was treated with
     Et isocyanoacetate in refluxing EtCOMe contg. Bu2Sn dilaurate to give
     glycinate III. III was active against Bacillus subtilus with a min.
     inhibitory concn. 10 ppm.
ST
     quinolinyl carbamate prepn urinary antimicrobial; glycine
     quinolinyloxycarbonyl prepn antimicrobial; bactericide quinolinyl
     carbamate; fungicide quinolinyl carbamate
     Bactericides, Disinfectants, and Antiseptics
ΙT
     Fungicides and Fungistats
        (quinolinyl carbamates)
IT
     Urinary tract
        (quinolinyl carbamates as antimicrobials for)
                                93775-51-0P 93775-52-1P
                                                           '93775-53-2P
ΙT
     19498-91-0P
                  51203-25-9P
                  93775-55-4P 93775-56-5P 93775-57-6P 93775-59-8DP, derivs.
     93775-54-3P 93775-55-4P
     93775-58-7P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); BIOL (Biological study); PREP (Preparation)
        (prepn. and antimicrobial activity of)
ΙT
     93775-51-0P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and hydrolysis and antimicrobial activity of)
                                 14种种、相邻、用类型14年
IT
     19642-75-2P
     RL: SPN (Synthetic preparation); PREP (Preparation)
                    (prepn. of)
                        7773-76-2
               521-74-4
                                   826-81-3 4008-48-4
IT
     130-16-5
                         1966年 - 1968年 1964年 1966年 11年
     RL: RCT (Reactant)
        (reaction of, with Et isocyanatoacetate)
IT
                       1 1 1 1 1 1 1 1 1
                                 (李克克·特斯科斯)。
     1943-83-5
                                  ind Antiberative
     RL: RCT (Reactant)
        (reaction of, with chlorohydroxyquinoline)
     2949-22-6
IT
     RL: RCT (Reactant)
        (reaction of, with hydroxyquinolines)
IT
     RL: BAC (Biological activity of effector, except adverse); SPN (Synthetic
     preparation); BIOL (Biological study); PREP (Preparation)
        (prepn. and antimicrobial activity of)
RN
     93775-55-4 HCAPLUS
     Glycine, N-[[(5-nitro-8-quinolinyl)oxy]carbonyl]-, ethyl ester (9CI)
CN
     INDEX NAME)
```

1,

the public of the factor of th

COPYRIGHT 2001 ACS ANSWER 21 OF 24 HCAPLUS L27 1977:568362 HCAPLUS AN DN 87:168362 E 3 . . . 11 Preparation and properties of some .alpha.-aza-amino-acid derivatives, ΤI their possible use in peptide synthesis Gray, C. J.; Ireson, J. C.; Parker, R. C. ΑU Dep. Chem., Univ. Birmingham, Birmingham, Engl. CS Tetrahedron (1977), 33(7), 739-43 CODEN: TETRAB SO DT Journal LA English CC 34-2 (Synthesis of Amino Acids and **Proteins**)

GΙ

Section cross-reference(s): 28

Azaglycine and azaphenylalanine derivs. were prepd. by acylation of AΒ acylhydrazides with alkyl and aryl chloroformates. E.g., Me3CO2CNHNH2 and AcNHNHCH2Ph with ClCO2Ph gave Me3CO2CNHNHCO2Ph (I) and AcNHN(CH2Ph)CO2Ph (II), resp. Esters of acetyl- and benzoylazaamino acids underwent rapid cyclization to oxadiazolones and were unsuitable for peptide synthesis. E.g., II gave the oxadiazolone III on treatment with NaOH or NH2OH or incubation at pH 7 at 37.degree for several days. Me3CO2CNHNHCON3, prepd. from I by sequential reactions with NH2NH2 and amyl nitrite was too unreactive for peptide synthesis Coupling reaction of 2,4-dinitrophenyloxycarbonylphenylalanine Et ester with BzNHNH2 and AcNHNH2CH2Ph gave benzoylazaglycyl#L-phenylalanine Et ester and acetylazaphenylalanyl-L-phenylalanine Et ester, resp. aza amino acid prepn cyclization; hydrazide acyl acylation; aza peptide; ST ring closure aza amino acid; oxadiazolone IT Amino acids, preparation RL: SPN (Synthetic preparation); PREP (Preparation) (aza-, prepn. of, by acylation of hydrazides) Ring closure and formation IT

IT Hydrazides
RL: RCT (Reactant)

高級 四代

்குதிற்க ர்த்ச நிறுக்க

1.34 · App population (App population)
 1.42 for any form (App population)
 1.46 form (App population)

(of aza amino acids, oxadiazolones by)

2 Post

```
(acyl, acylation of, aza amino acids by)
                                  1. $45 月1
    Peptides, preparation
IT
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (aza-, prepn. of, by reaction of peptides with acylhydrazides)
               870-46-2! †\1215-52<del>-</del>7\!\\7151-53-3\!\\53370-84-6
IT
     613-94-5
    RL: RCT (Reactant)
        (acylation of)
ΙT
     3081-24-1
    RL: RCT (Reactant)
        (dinitrophenyloxycarbonylation of)
IT
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and azidation of)
                                   प्रिक्षा पति । जप्रक
ΙT
     64512-92-1P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and condensation reactions of, with benzoylhydrazide and
       acetylbenzylhydrazine)
                                64512-88-5P
IT
     53370-82-4P
                  53370-85-7P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and cyclization of)
     64512-91-0P
IT
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and reaction of, with benzylamine)
    1199-02-6P
                               27643-12-5P 52414-76-3P
                                                           53370-83-5P
IT
                 15081-44-4P
                                64512 - 81 - 8P^{1} = 64512 + 82 - 9P
                  57699-88-4P
                                                            64512-83-0P
     57699-63-5P
                                64512-86-3P 64512-87-4P
                                                            64512-93-2P
    64512-84-1P
                  64512-85-2P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
     64512-89-6P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn., deprotection, and reactions of, with ammonia and hydrazine)
IT
     64512-92-1P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and condensation reactions of, with benzoylhydrazide and
        acetylbenzylhydrazine)
                                  left fishell with min H
     64512-92-1 HCAPLUS
RN
     CN
     INDEX NAME)
Absolute stereochemistry
        NO2
```

```
COPYRIGHT 2001 ACS
L27
    ANSWER 22 OF 24 | HCAPLUS
    1975:4556 HCAPLUS
ΑN
                                  翻入时间 随时间身
DN
    82:4556
    Hydrazine compounds as hetero-components in peptides. XIX. Eledoisin
TI
    peptides containing the N2-methylcarbazoyl radical (azaalanine)
    Niedrich, Hartmut; Oehme, Christa; Bergmann, Jutta
ΑU
    Zentralinst. Molekularbiol., DAW, Berlin, E. Ger.
CS
    J. Prakt. Chem. (1974), 316(5), 741-9
SO
    CODEN: JPCEAO
```

```
DT
           Journal
LΑ
           German
CC
           34-3 (Synthesis of Amino Acids,
           and Pr teins)
           Acylation, sapon., and peptide coupling of 2+azaalanine peptides were
AΒ
           studied with H2NNMeCONHCHPhCO2Me (Azala-Phe+OMe) (T). 6-Azala-eledoisin
           sequence 6-11 (II) and 5-Ala-6-Azala-eledoisin 5-11 (III) were prepd. by
           coupling of I with Ile-Gly-Leu-Met-NH2 in the presence of
           hydroxysuccinimide and dicyclohexylcarbodiimide. III and III had only the
           biol. activity of the 7-11 sequence, i.e. .apprx.18 of the 6-11 sequence.
           azaalanine peptide eledoisin
ST
ΙT
           Peptides, preparation
           RL: PREP (Preparation)
                  (azaalanine-contg., eledoisin-related)
IT
           3069-69-0
           RL: RCT (Reactant)
                  (methylation of)
IT
           40203-94-9
           RL: RCT (Reactant)
                  (peptide coupling reaction with azaalanine derivs.)
ΙT
           2577-90-4
           RL: RCT (Reactant)
                  RCT (Reactant)
(peptide coupling reactions of)
                                      2280-71+94日 2304-96-36日2592-19-0
ΙT
           1142-20-7
                                                                                                                       15761-38-3
           RL: RCT (Reactant)
                  (peptide coupling reactions with azaalanine deriv.)
                                          54601-22-8P
                                                                          54601+24#0P 1 54601+25#1P
IT
           54601-18-2P
                                                                           54601-28-4P
           54601-27-3P
           RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
                  (prepn. and peptide coupling reactions of)
                                                                          54601-23-9P
                                                                                                          54601-29-5P
                                          54601-20-6P
IT
           54601-19-3P
                                                                                                           The state of the s
                                           54601-32-0P
                                                                           54601-33-1P
           54601-31-9P
           RL: SPN (Synthetic preparation); PREP (Preparation)
                  (prepn. of)
           60-34-4
IT
           RL: RCT (Reactant)
                  (reaction of, with amino acids)
IT
           2483-49-0 54601-21-7;
           RL: RCT (Reactant)
                  (reaction of, with methylhydrazine)
IT
           54601-21-7
           RL: RCT (Reactant)
                  (reaction of, with methylhydrazine)
RN
           54601-21-7 HCAPLUS: 18-41
CN
           L-Phenylalanine, N-[(4-nitrophenoxy) carbonyl]-, methyl ester (9CI)
           INDEX NAME)
                                                                                        :世界東
Absolute stereochemistry
```

L27 ANSWER 23 OF 24 HCAPLUS COPYRIGHT 2001 ACS

[南阳學學]第6報第145年7年上

```
1970:425854
                  HCAPLUS
AN
DN
     73:25854
     Cyclizations via phenylthio- and phenoxycarbonylamino intermediates. II.
ΤI
     Formation of azasuccinic anhydrides and their polycondensation products.
     Pyridiniocarbonylamino intermediates
     Baudet, Pierre; Otten; Cl.; Rao, D.
AU
CS
     Lab. Chim. Org., Univ. Geneve, Geneva, Switz.
     Helv. Chim. Acta (1970), 53(4), 859-69
SO
     CODEN: HCACAV
DT
     Journal
                                    4:4-75
     French
LА
     34 (Synthesis of Amino Acids, Peptides, and
CC
     The N-(p-nitrophenoxycarbonyl) derivs of glycine, DL-alanine and
AΒ
     DL-leucine were transformed by pyridine into azasuccinic,
     3-methyl-2-azasuccinic and 3-isobutyl-2-azasuccinic anhydride, resp., by
     way of N-carbamoylpyridinium cation intermediates. The above azasuccinic
     anhydrides polycondensed in the presence of pyridine yielding the corresponding polyglycine, poly-DL-alanine and poly-DL-leucine, resp.
     Both cyclization and polycondensation were also catalyzed by
     .gamma.-collidine, but at a low rate. N-(p-Nitrophenoxycarbonyl)glycine
     reacted with lysozyme in the presence of pyridine. Several glycine
     residues were introduced into the enzyme which was rendered insol. and
     partially inactivated.
                                   azasuccinic anhydridės; anhydridės azasuccinić; polyamino acids; lysozyme
ST
     glycyl; glycyllysozyme
                                   1917年群
IT
     Pyridinium compounds
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (carbonyl derivs. of amino acids, intermediates in oxazolidinedione
        prepn.)
     2,5-Oxazolidinedione, derivs.
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of, from phenoxycarbonyl amino acids and pyridine)
                  2185-00-4P 4289-99-0P 21639-02-1P 25281-63-4P
IT
     1192-73-0P
                                  25988=64-1P 26283-00-1P
                                                               26334-33-8P
                   25734-27-4P
     25718-94-9P
     26952-07-8P 27317-15-3P 27317-16-4P 27317-17-5P 27317-18-6P 27317-19-7P
     27317-18-6P 27317-19-7P
     RL: SPN (Synthetic preparation) PREP (Preparation) (prepn. of)
21639-02-1P 27317-15-3P 27317-16-4P
27317-18-6P 27317-19-7P
IT
     RL: SPN (Synthetic preparation) PREP (Preparation)
                        preparation) 7, rape A
        (prepn. of)
     21639-02-1 HCAPLUS
RN
     Glycine, N-carboxy-, N-(p-nitrophenyl) ester (8CI) (C
                                                         (CA INDEX NAME)
CN
                                  通过的第三人称单数
                                    hir approved to the first of the
                                    PARTHURFACED FOLLS IN
02N
                         1 Po 22
                                    27317-15-3 HCAPLUS
RN
     Alanine, N-carboxy-, ethyl N-(p-nitrophenyl) ester, DL- (8CI) (CA INDEX
CN
     NAME)
                                   information appropriate
```

的物质分别 拉拉克克克拉

RNHCAPLUS

N-(p-nitrophenyl) ester, DL- (8CI) (CA INDEX NAME) CN Alanine, N-carboxy-,

RN27317-18-6 HCAPLUS

Leucine, N-carboxy-, ethyl N-(p-nitrophenyl) ester, DL- (8CI) (CA INDEX CN NAME)

RN27317-19-7 HCAPLUS

Leucine, N-carboxy-, N-(p-nitrophenyl) ester, DL- (8CI) CN

ANSWER 24 OF 24 HCAPLUS COPYRIGHT 2001 ACS L27

AN 1967:18854 HCAPLUS

DN 66:18854

持持續的 Peptide syntheses. XXXII. .gamma.-Hydroxyisocaproyl and ΤI 3-nitrophenoxycarbonyl moieties as protecting groups

Wieland, Theodor; Lamperstorfer, Ch.; Birr, Christian ΑU

Univ. Frankfurt/M., Frankfurt/M., Ger. CS

Makromol. Chem. (1966), 92, 277-86 SO

CODEN: MACEAK

DT Journal

German LΑ

CC 34 (Synthesis of Amino Acids, Peptides, and Proteins)

cf. CA 64, 14272g. The title protecting groups were found to react readily with various amino acids, and to be removed quant. by CF3CO2H treatment and irradiation with uv light (lambda. >290 m.mu.), resp. Thus, a soln. of 15 g glycine (I) in 100 ml. 2N NaOH was evapd. to dryness in vacuo and the residue dried in a desiccator. The resulting glycine Na salt (9.7 g.) was pulverized and dissolved in 100 g. molten imidazole (II) on a boiling water bath. Isocaprolactone (16 g.) prepd. according to Stevens and Tarbell (CA 50, 937e) was added, the mixt. was heated 1 hr., cooled, and washed several times with Me2CO to give 43% N-(.gamma.-hydroxyisocaproyl)glycine (III) Na salt, mixed with a little I Na salt. The crude salt (4.2 g.) was dissolved in 10 ml. H2O, cooled in ice, acidified to pH 2.6 with 0.1N H2SO4, satd. with NaCl, and extd. with five 10 ml. vols. AcEt. The org. exts. were evapd. to a vol. of 20 ml. and treated with petr. ether to give 1.6 g. III, m. 106.degree. (AcoEt-petr. ether). In a similar fashion, N-(.gamma.-hydroxy-isocaproyl)-L-leucine (IV), which sintered at 190.degree. (decompn.), was prepd. in 10% yield. A mixt. of 600 mg. III and L-leucine Me ester hydrochloride was converted by the anhydride method of Determann, et al. (CA 57, 9948c) to N-(.gamma.-hydroxyisocaproyl) -glycyl-L-leucine Me ester in 36% yield. In a similar manner, 390 mg. IV and L-tyrosine benzyl ester tosylate were converted to 50% N-(.gamma.-hydroxyisocaproyl)-L-leucyl-L-tyrosine benzyl ester (V), m. 154-5.degree.. Total hydrolysis of V gave equal amts. leucine (VI) and tyrosine. The gamma hydroxyisocaproyl group was found most difficult to remove in the case of V; 10 mg. of which was dissolved in 0.2 ml. 50% CF3CO2H soln., kept 12 hrs. at room temp., and evapd. in a desiccator over concd. H2SO4 and KOH. The residue was shown by electrophoresis to be L-leucyl-L-tyrosine benzyl ester. For the prepn. of 3-nitrophenoxycarbonyl derivs : 38 g : COC12 was dissolved in 160 ml. ice-cold abs. C6H6. The soln. was stirred, treated dropwise during 2 hrs. with a soln. of 30 g m-O2NC6H4OH (VII) and 36.5 g. PhNMe2 in AcOEt at 5-10.degree., and stirred 12 hrs. at room temp. The mixt. was extd. 3 times with 100-ml. vols. N HCl, washed with 100 ml. H2O, and the org. phase was evapd. to dryness in vacuo to give 74% 3-02NC6H402CCl (VIII), b13 150-1.degree.. A mixt. of 6.5 g. VI, 4 g. MgO, 100 ml. H2O, and 30 ml. Et20 was cooled in ice, stirred vigorously, and treated during 30 min. with a mixt. of 10 g. VIII dild. with ancequal vol. Et20. The mixt. was acidified with 30 ml. concd. H2SO4 and extd. with three 30-ml. vols. The combined org. exts. were washed 3 times with 20-ml. vols. 2N HCl and 5 times with 20-ml. vols H20, dried, and evapd. to dryness in vacuo at 30.degree. to give 62% 3 introphenoxycarbonyl-L-leucine, mixed with some VII. In a similar fashion, L-phenylalanine (IX) was converted to 52% 3-nitrophenoxycarbonyl deriv. (X) , m. 118.degree. (AcoEt-petr. ether). A soln. of 3.3 g. X and 0.89 g. NH2CH2CO2Me in 50 ml. abs. tetrahydrofuran (XI) was cooled to -15 degree . Stirred, and treated with 0.92 ml. POC13, followed at once by 2.8 ml. Et3N. The mixt. was stirred 1 hr. at -15.degree., treated with 20 ml. H20, and concd. in vacuo. The residue was extd. with three 10-ml. vols. AcoEt, which exts. were combined and washed with 10 ml. H2O, 10 ml. 5% aq. NaHCO3, and three 10-ml. vols. H2O, dried, and evapd to dryness in vacuo at 30 degree. to give 32.5 g. 3-nitrophenoxycarbonyl-L-phenylalanylglycine Me ester, m. 162.degree. (AcOEt-petr. ether and MeOH-H2O), still mixed with VII. Et3N (1.43 ml.) and then 0.95 ml. ClCO2Et was added with shaking to a soln. of 3.3 g. X in 20 ml. XI, and the mixt. cooled to 15 degree . After 8 min., the mixt. was warmed to 0.degree., treated with a soln. of 2.7 g. L-alanylglycine benzyl ester hydrochloride in 30 ml. XI and 20 ml. H2O (mixed just before addn. with 1.43 ml. Et3N), and removed from the cooling bath and shaken until there was a strong evolution of CO2. The soln was evapd. in vacuo at 40.degree., and the residue mixed with 60 ml. AcoEt, which phase was washed with two 10-ml. vols. each of Nº HCl and H2O, two vols. 5% aq. NaHCO3, and five 10-ml. vols. H2O, dried, and evapd. in vacuo at

海线 经营工工程 医自己性

AB

30.degree. to give 75% 3-nitrophenoxycarbonyl-L-phenylalanylglycyl-L-alanine benzyl ester, m. 180-1 degree. (MeOH-H2O). The photolysis of X was performed by irradiating a soln. of 0.98 g. X in 200 ml. 25% aq. XI with a water-cooled high-pressure-Hg lamp (Hanau Q 81). Paper electrophoresis showed hydrolysis to be complete after 4 hrs., and 89% IX was recovered. Expts. to cleave the 3-nitrophenoxy-carbonyl group with aq. Et3N led to hydantoin formation. 18 references.

NITROPHENOXYCARBONYL PEPTIDE PROTECTIVE GROUP; HYDROXYISOCAPROYL PEPTIDE PROTECTING GROUPS HYDROXYISOCAPROYL; NITROPHENOXYCARBONYL PEPTIDE PROTECTIVE GROUP; PEPTIDE PROTECTING GROUPS HYDROXYISOCAPROYL; HYDROXYISOCAPROYL PEPTIDE PROTECTING GROUPS

IT Peptides, preparation

RL: PREP (Preparation)

(4-hydroxy-4-methylvaleryl group and (m-nitrophenoxy) carbonyl group as protective groups for amino group in)

IT Amino group

(4-hydroxy-4-methylvaleryl group and (m-nitrophenoxy) carbonyl group as protective groups for amino group in peptide prepn.)

IT (m-Nitrophenoxy) carbonyl group

4-Hydroxy-4-methylvaleryl group

(as protective group for amino group in peptide prepn.)

14235-02-0P 14235-03-1P 14235-04-2P 14235-05-3P 14235-06-4P 14235-07-5P 14235-13-3P 14317-60-3P 14235-05-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

IT 14235-06-4P 14235-13-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

RN 14235-06-4 HCAPLUS

CN Alanine, N-carboxy-3-phenyl-, N-m-nitrophenyl) ester, L- (8CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 14235-13-3 HCAPLUS

CN Leucine, N-carboxy-, N-(m-nitrophenyl) ester, L- (8CI) (CA INDEX NAME)

Absolute stereochemistry.